# Exhibit 3

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(54) Title: MONOCLONAL ANTIBODIES WITH RED	UCED	IMMUNOGENICITY
(57) Abstract		
Antibodies having reduced immunogenicity and me	thods f	or making them are disclosed.
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### MONOCLONAL ANTIBODIES WITH REDUCED IMMUNOGENICITY

This application claims the benefit of U.S. Provisional Application No. 60/083,367, filed April 28, 1998.

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### Field of the Invention

This invention relates to monoclonal antibodies (mAbs) having reduced immunogenicity in humans.

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### Background of the Invention

Many potentially therapeutic mAbs are first generated in a murine hybridoma system for reasons of speed and simplicity. Non-human mAbs contain substantial stretches of amino acid sequences that will be immunogenic when injected into a human patient. It is well known that after injection of a foreign antibody, such as a murine antibody, a patient can have a strong human anti-mouse antibody (HAMA) response that essentially eliminates the antibody's therapeutic utility after the initial treatment as well as the utility of any other subsequently administered murine antibody.

Humanization techniques are well known for producing mAbs which exhibit reduced immunogenicity in humans while retaining the binding affinity of the original non-human parental mAb. See, e.g., those disclosed in U.S. Patent Nos. 5,585,089; 5,693,761; 5,693,762; and 5,225,539.

In general, these methods depend on replacing human variable heavy and light region complementarity determining regions (CDRs) with antigen specific non-human CDRs, a process known as CDR grafting. It is also well known that in CDR grafting experiments the retention of the original antigen binding affinity is enhanced and in many cases depends on choosing human acceptor framework regions that most closely match the corresponding frameworks of the CDR donor antibody.

However, since the human genome contains a limited repertoire of heavy and light chain framework regions, these methods suffer from the limitation of available human acceptor frameworks. This restriction in acceptor framework repertoire necessarily can limit the degree of match between the non-human donor and the human acceptor antibody. Thus,

CDR grafting methods are limited by the known available repertoire of human VH and VL framework regions. Clearly, a need exists for an expanded range of acceptor V regions.

### Summary of the Invention

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One aspect of the present invention is an antibody comprising donor CDRs derived from an antigen-specific donor antibody of a non-human species and acceptor framework residues derived from a non-human primate.

Another aspect of the invention is a method for making an antibody having reduced immunogenicity in humans comprising grafting CDRs from antigen-specific non-human antibodies onto homologous non-human primate acceptor frameworks.

Another aspect of the invention is a chimpanzee VH acceptor framework I, II and III comprising an amino acid sequence as set forth in SEQ ID NOs: 10, 11, 12, 13, 14, 15, 16, 17 or 18.

Another aspect of the invention is a chimpanzee VH acceptor framework IV comprising an amino acid sequence as set forth in SEQ ID NOs: 81, 82, 83, 84 or 85.

Another aspect of the invention is a chimpanzee VK acceptor framework I, II and III comprising an amino acid sequence as set forth in SEQ ID NOs: 28, 29, 30, 31, 32, 33, 34, 35 or 36.

Another aspect of the invention is a chimpanzee  $V\kappa$  acceptor framework IV comprising an amino acid sequence as set forth in SEQ ID NOs: 86 or 87.

Another aspect of the invention is a cynomolgus VH acceptor framework I, II and III comprising an amino acid sequence as set forth in SEQ ID NOs: 45, 46, 47, 48, 49, 50, 51 or 52.

Another aspect of the invention is a cynomolgus VH acceptor framework IV comprising an amino acid sequence as set forth in SEQ ID NOs: 88, 89, 90, 91, 92 or 93.

Another aspect of the invention is a cynomolgus VK acceptor framework I, II and III comprising an amino acid sequence as set forth in SEQ ID NOs: 59, 60, 61, 62, 63 or 64.

Another aspect of the invention is a cynomolgus  $V\kappa$  acceptor framework IV comprising an amino acid sequence as set forth in SEQ ID NOs: 94, 95 or 96.

Yet another aspect of the invention is an isolated nucleic acid molecule encoding the amino acid sequence of SEQ ID NOs: 10, 11, 12, 13, 14, 15, 16, 17, 18, 28, 29, 30, 31, 32, 33, 34, 35 or 36.

Yet another aspect of the invention is an isolated nucleic acid molecule encoding the amino acid sequence of SEQ ID NOs: 81, 82, 83, 84, 85, 86 or 87.

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Yet another aspect of the invention is an isolated nucleic acid molecule encoding the amino acid sequence of SEQ ID NOs: 45, 46, 47, 48, 49, 50, 51, 52, 59, 60, 61, 62, 63 or 64.

Yet another aspect of the invention is an isolated nucleic acid molecule encoding the amino acid sequence of SEQ ID NOs: 88, 89, 90, 91, 92, 93, 94, 95 or 96.

### Brief Description of the Drawings

Figure 1 is an amino acid sequence of the engineered 4A6 VL region. Asterisks above the 4A6 sequence indicate the 4A6 framework residues retained in the engineered molecule. Bold and italicized letters indicate the CDRs.

Figure 2 is an amino acid sequence of the engineered 4A6 VH region. Asterisks above the 4A6 sequence indicate the 4A6 framework residues retained in the engineered molecule. Bold and italicized letters indicate the CDRs.

Figure 3 is an amino acid sequence alignment comparing the murine antibody B9Vk with the closest matching chimpanzee Vk and selected Jk sequences. The CDR regions are indicated by bold and italicized letters. Gaps are indicated by dots. The numbering convention is from Kabat et al., infra.

Figure 4 is an amino acid sequence alignment comparing the murine antibody B9VH with the closest matching chimpanzee VH and selected JH sequences. The CDR regions are indicated by bold and italicized letters. Gaps are indicated by dots. Asterisks indicate framework residues that are predicted to interact with CDRs and affect antigen binding affinity. The numbering convention is from Kabat et al., infra.

Figure 5 is an amino acid sequence alignment comparing the murine antibody 3G9Vk with the closest matching chimpanzee Vk and selected Jk sequences. The CDR regions are indicated by bold and italicized letters. Gaps are indicated by dots. Asterisks indicate framework residues that are predicted to interact with CDRs and affect antigen binding affinity. The numbering convention is from Kabat et al., infra.

Figure 6 is an amino acid sequence alignment comparing the murine antibody 3G9VH with the closest matching chimpanzee VH and selected JH sequences. The CDR regions are indicated by bold and italicized letters. Gaps are indicated by dots. Asterisks indicate framework residues that are predicted to interact with CDRs and affect antigen binding affinity. The numbering convention is from Kabat et al., infra.

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### Detailed Description of the Invention

All publications, including but not limited to patents and patent applications, cited in this specification are herein incorporated by reference as though fully set forth.

The molecular genetic aspects of antibody structure have been reviewed by S. Tonegawa in Nature 302:575-581 (1983). Briefly, antibodies are heterodimers comprised of at least two heavy and two light chains. The N-terminal domain of each heavy and light chain, termed VH and VL, respectively, fold together to form the antigen combining site. On the genetic level, the VL domain is encoded by two different gene segments, termed VK or Vl, and JK or Jl that join together to form one continuous VL region. Similarly, the VH domain is encoded by three gene segments, VH, DH, and JH, that join together to form one continuous VH region. Thus different VL and VH regions may be encoded by different combinations of VK or Vl, Jk or Jl and VH, DH, and JH. This combinatorial diversity is in part the means by which the immune response generates the myriad diversity of different antibody molecules and their associated antigen specificities.

On the protein level, each heavy and light V region domain may be further divided into three CDRs. Three heavy

and three light chain CDRs fold together to form the antigen binding surface and part of the underlying support structures that are required to maintain the exact three-dimensional structure of the antigen combining site. Flanking each CDR are framework regions that in most cases do not directly interact with the specific antigen, but rather serve to form the scaffold which supports the antigen binding properties of the CDRs. Each heavy and light chain has four framework regions, three derived from the VH or VL gene segment, the fourth is derived from the JH, JK, or Jl gene segment. the order of frameworks and CDRs from the N- terminus is framework I, CDRI, framework II, CDRII, framework III, CDRIII, framework IV. On the genetic level, all of framework I through Framework III is encoded by the V region gene segment; CDRIII is encoded jointly by both the V region and J region gene segments; framework IV is encoded entirely from the J gene segment.

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As used herein, "antibodies" refers to immunoglobulins and immunoglobulin fragments lacking all or part of an immunoglobulin constant region, e.g., Fv, Fab, Fab' or  $F(ab')_2$  and the like.

The term "donor antibody" refers to a monoclonal or recombinant antibody which contributes the nucleic acid sequences of its variable regions, CDRs or other functional fragments or analogs thereof to an engineered antibody, so as to provide the engineered antibody coding region and resulting expressed engineered antibody with the antigenic specificity and neutralizing activity characteristic of the donor antibody.

The term "acceptor antibody" refers to monoclonal or recombinant antibodies heterologous to the donor antibody, which contributes all, or a portion, of the nucleic acid sequences encoding its heavy and/or light chain framework regions and/or its heavy and/or light chain constant regions or V region subfamily consensus sequences to the engineered antibody.

A "functional fragment" is a partial heavy or light chain variable sequence (e.g., minor deletions at the amino or carboxy terminus of the immunoglobulin variable region)

which retains the same antigen binding specificity and affinity as the antibody from which the fragment was derived.

An "analog" is an amino acid sequence modified by at least one amino acid, wherein said modification can be chemical or a substitution, which modification permits the amino acid sequence to retain the biological characteristics, e.g., antigen specificity and high affinity, of the unmodified sequence.

Methods are provided for making engineered antibodies with reduced immunogenicity in humans and primates from non-10 human antibodies. CDRs from antigen-specific non-human antibodies, typically of rodent origin, are grafted onto homologous non-human primate acceptor frameworks. Preferably, the non-human primate acceptor frameworks are from Old World apes. Most preferably, the Old World ape 15 acceptor framework is from Pan troglodytes, Pan paniscus or Gorilla gorilla. Particularly preferred is the chimpanzee Pan troglodytes. Also preferred are Old World monkey acceptor frameworks. Most preferably, the Old World monkey acceptor frameworks are from the genus Macaca. Particularly 20 preferred is the cynomolgus monkey Macaca cynomolgus.

Particularly preferred chimpanzee (Pan troglodytes) heavy chain variable region frameworks (VH) are CPVH41-12 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 10 and the framework IV amino acid sequence shown in SEQ ID NO: 83; CPVH41-1 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 11 and the framework IV amino acid sequence shown in SEQ ID NO: 85; CPVH41-4 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 12; CPVH41-7 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 13; CPVH41-8 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 14, CPVH41-9 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 15 and the framework IV amino acid sequence shown in SEQ ID NO: 81; CPVH41-10 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 16 and the framework IV amino acid sequence shown in SEQ ID NO: 82; CPVH41-18 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 17; and CPVH41-19 having the framework I, II and III

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amino acid sequence shown in SEQ ID NO: 18 and the framework IV amino acid sequence shown in SEQ ID NO: 84.

Particularly preferred chimpanzee (Pan troglodytes) light chain kappa variable region frameworks (VK) are CPVK46-1 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 28; CPVK46-3 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 29; CPVK46-4 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 30; CPVK46-5 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 31; CPVK46-6 having the 10 framework I, II and III amino acid sequence shown in SEQ ID NO: 32 and the framework IV amino acid sequence shown in SEQ ID NO: 86; CPVK46-7 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 33 and the framework IV amino acid sequence shown in SEQ ID NO: 87; CPVK46-8 having 15 the framework I, II and III amino acid sequence shown in SEQ ID NO: 34; CPVK46-11 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 35; and CPVK46-14 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 36. 20

Particularly preferred cynomolgus (Macaca cynomolgus) heavy chain variable region frameworks (VH) are CYVH2-1 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 45 and the framework IV amino acid sequence shown in SEQ ID NO: 88; CYVH2-3 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 46 and the framework IV amino acid sequence shown in SEQ ID NO: 89; CYVH2-4 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 47 and the framework IV amino acid sequence shown in SEQ ID NO: 90; CYVH2-5 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 48 and the framework IV amino acid sequence shown in SEQ ID NO: 93; CYVH2-6 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 49 and the framework IV amino acid sequence shown in SEQ ID NO: 91; CYVH2-7 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 50; CYVH2-8 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 51; and CYVH2-10 having the

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framework I, II and III amino acid sequence shown in SEQ ID NO: 52 and the framework IV amino acid sequence shown in SEQ ID NO: 92.

Particularly preferred cynomolgus (Macaca cynomolgus) light chain kappa variable region frameworks (VK) are CYVK4-2 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 59; CYVK4-3 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 60 and the framework IV amino acid sequence shown in SEQ ID NO: 94; CYVK4-5 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 61; CYVK4-6 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 62 and the framework IV amino acid sequence shown in SEQ ID NO: 95; CYVK4-10 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 63; and CYVK4-11 having the framework I, II and III amino acid sequence shown in SEQ ID NO: 64 and the framework IV amino acid sequence shown in SEQ ID NO: 64 and the framework IV amino acid sequence shown in SEQ ID NO: 96.

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Isolated nucleic acid molecules encoding the chimpanzee VH and Vk acceptor framework I, II and III amino acid sequences of SEQ ID NOs: 10, 11, 12, 13, 14, 15, 16, 17, 18, 28, 29, 30, 31, 32, 33, 34, 35 or 36 and the framework IV amino acid sequences of SEQ ID NOs: 81, 82, 83, 84,85, 86 or 87 are also part of the present invention. Further, isolated nucleic acid molecules encoding the cynomolgus VH and Vk acceptor framework I, II and III amino acid sequences of SEQ ID NOs: 45, 46, 47, 48, 49, 50, 51, 52, 59, 60, 61, 62, 63 or 64 and the framework IV amino acid sequences of SEQ ID NOs: 88, 89, 90, 91, 92, 93, 94, 95 or 96 are also part of the present invention. Nucleic acid sequences encoding functional fragments or analogs of the VH and Vk acceptor framework amino acid sequences are also part of the present invention.

In addition to isolated nucleic acid sequences encoding VH and Vk acceptor frameworks described herein, nucleic acid sequences complementary to these framework regions are also encompassed by the present invention. Useful DNA sequences include those sequences which hybridize under stringent hybridization conditions to the DNA sequences. See, T.

Maniatis et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory (1982), pp. 387-389. An example of one such stringent hybridization condition is hybridization at 4XSSC at 65°C, followed by a washing in 0.1XSSC at 65°C for one hour. Alternatively, an exemplary stringent hybridization condition is 50% formamide, 4XSSC at 42°C. Preferably, these hybridizing DNA sequences are at least about 18 nucleotides in length.

Suitable frameworks are selected by computer homology searching among members of a database of Old World ape or monkey VH and VL regions. The framework portions of primate antibodies are useful as components of therapeutic antibodies. Moreover, primate antibody frameworks will be tolerated when used in the treatment of humans due to the close sequence homology between the genes of the primates and humans. Thus, the present invention provides for the grafting of CDRs from an antigen specific non-human donor antibody to acceptor V regions derived from non-human primate species.

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The antigen specificity and binding kinetics of the donor antibody, which may be of rodent or any other non-human origin, are best preserved by selecting primate acceptor V regions that are determined by computer homology searching to be most similar to the donor antibody. Alternatively, the acceptor antibody may be a consensus sequence generated from primate V region subfamilies, or portions thereof, displaying the highest homology to the donor antibody.

The resulting engineered constructs, in which the donor CDRs are grafted onto primate acceptor frameworks, are subsequently refined by analysis of three-dimensional models based on known antibody crystal structures as found, e.g., in the Protein Data Bank, http://www.pdb.bnl.gov/pdb-bin/pdbmain. Alternatively, computer generated three-dimensional models of the donor antibody may be computed by means of commercially available software such as "AbM" (Oxford Molecular, Oxford, UK).

Structural analysis of these models may reveal donor framework residues that are CDR-contacting residues and that are seen to be important in the presentation of CDR loops,

and therefore binding avidity. A CDR-contacting residue is one which is seen in three-dimensional models to come within the van der Waals radius of a CDR residue, or could interact with a CDR residue via a salt bridge or by hydrophobic interaction. Such donor framework (CDR-contacting) residues may be retained in the engineered construct.

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The modeling experiments can also reveal which framework residues are largely exposed to the solvent environment. The engineered constructs may be further improved by substituting some or all of these solvent-accessible amino acid residues with those found at the same position among human V regions most homologous to the engineered construct as disclosed in U.S. Patent No. 5,639,641.

The engineered V regions are then joined to one or more different human or Old World ape constant regions depending on the desired secondary immune functions such as complement fixation or Fc receptor binding. Human constant regions can be selected from human immunoglobulin classes and isotypes, such as IgG (subtypes 1 through 4), IgM, IgA, and IgE. An IgG4 subtype variant containing the mutations S228P and L235E (PE mutation) in the heavy chain constant region which results in reduced effector function can also be selected. See U.S. Patent Nos. 5,624,821 and 5,648,260.

The complete heavy and light chain genes are transferred to suitable expression vectors and co-expressed in the appropriate host cells such as chinese hamster ovary, COS or myeloma cells. The resulting engineered antibody is expected to be of substantially reduced immunogenicity when administered to humans, and to retain full binding affinity for antigen.

Acceptor V regions can be isolated specifically for each donor V region by directed PCR methodology where a non-human primate cDNA library is surveyed for acceptor frameworks most similar to the donor antibody. Oligonucleotide PCR primers homologous to the donor antibody framework I (paired with Cregion 3' PCR primers) are used to direct PCR amplification of a non-human primate, e.g., chimpanzee lymphocyte cDNA library. This would select for V-regions with framework I regions similar to the donor antibody, and sequence analysis of the obtained clones would reveal the associated framework

II and III (and IV) sequences. 3' PCR primers would then be designed based on the knowledge of the non-human primate framework III sequences thus obtained, and used to direct PCR amplification of the original cDNA library together with a vector-specific 5' PCR primer. cDNA clones obtained from the second round of PCR amplification would have framework I and III sequences most similar to the donor antibody, and the framework II sequences would display a similar degree of sequence homology.

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The present invention will now be described with reference to the following specific, non-limiting examples.

### Example 1

## 15 Random cDNA Cloning and Sequence Analysis of Chimpanzee VH Regions

Five ml of peripheral blood was collected and pooled from three chimpanzees (Pan troglodytes) and peripheral blood mononuclear cells were isolated by standard density centrifugation methods. These cells, which include antibody producing lymphocytes, were dissolved in TRIzol reagent (GIBCO, Gaithersburg, MD, USA) and total RNA was recovered from this material by solvent extraction and precipitation according to the manufacturer's specifications.

Chimpanzee heavy chain V regions were cloned from the total RNA using Marathon RACE methodology (Clontech, Palo Alto, CA, USA) following exactly the manufacturer's protocol using 3' Cg1 gene specific primers. After RACE PCR amplification, DNA bands of the expected size were excised from agarose gels, the DNA was purified and cloned into a plasmid vector. Although this cDNA library contains many distinct heavy chain V region clones, nine were selected randomly for sequence analysis. Complete nucleic acid sequences and predicted protein sequences of the chimpanzee VH cDNA clones 41-12, 41-1, 41-4, 41-7, 41-8, 41-9, 41-10, 41-18 and 41-19 are shown in SEQ ID NOs: 1, 2, 3, 4, 5, 6, 7, 8 and 9, respectively. The amino acid sequences of the region from the first amino acid of the mature VH region to the second conserved cysteine residue at position 92, adjacent to CDR III of these clones, namely, CPVH41-12,

CPVH41-1, CPVH41-4, CPVH41-7, CPVH41-8, CPVH41-9, CPVH41-10, CPVH41-18 and CPVH41-19 are shown in SEQ ID NOs: 10, 11, 12, 13, 14, 15, 16, 17 and 18, respectively. The amino acid sequence of the region encoding framework IV of these clones for CPVH41-9, CPVH41-10, CPVH41-12, CPVH41-19 and CPVH 41-1 are shown in SEQ ID NOs: 81, 82, 83, 84 and 85, respectively.

The chimpanzee VH amino acid sequences from the mature N-terminus and the second conserved cysteine residue at position 92, adjacent to CDRIII, were used as query sequences in computer homology searching of the Kabat database of Sequences of Proteins of Immunological Interest (ftp://ncbi.nlm.nih.gov/repository/kabat/) The results of this analysis are shown in Table 1.

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In each case, the closest match was with a human VH region, displaying between 76% (41-1/HHC20G) and 94% (41-10/HHC20Y) sequence identity at the amino acid level.

Matches were found for each of the three major human VH subgroups, indicating that the chimpanzee VH repertoire includes at least some members homologous to each of the major human subgroups. The human subgroup homology is presented in Table 1.

25			<b>Table 1</b> Overall Amino	
	Clone	Closest Match	Acid Homology	VH Subgroup Match
	41-4	HHC10X	88%	I
	41-9	HHC10Y	92	I
	41-18	HHC10D	84	I
30	41-1	HHC20G	76	II
	41-10	HHC20Y	94	II
	41-12	HHC20C	83	II
	41-7	HHC30T	80	III
	41-8	HHC30T	79	III
35	41-19	HHC305	82	III

The results show that the overall sequence identity between the chimpanzee and human VH regions ranged between 76 and 95% with a mean identity of 84%. Based on this observation, further sampling of the chimpanzee random VH library will likely provide a substantially greater diversity of VH sequences from which to choose optimum acceptor frameworks for each particular donor VH region.

#### Example 2

## Random cDNA Cloning and Sequence Analysis of Chimpanzee VK Regions

Chimpanzee light chain VK regions were cloned from the total RNA using Marathon RACE methodology (Clontech, Palo Alto, CA, USA) following exactly the manufacturer's protocol and Ck 3' gene specific primers. After RACE PCR amplification, DNA bands of the expected size were excised from agarose gels, the DNA was purified and cloned into a plasmid vector. Although this cDNA library contains many 1.0 distinct light chain VK region clones, nine were selected randomly for sequence analysis. Complete nucleic acid sequences and predicted protein sequences of the chimpanzee VK cDNA clones 46-1, 46-3, 46-4, 46-5, 46-6, 46-7, 46-8, 46-11 and 46-14 are shown in SEQ ID NOs: 19, 20, 21, 22, 23, 24, 15 25, 26 and 27, respectively. The amino acid sequences of the region from the first amino acid of the mature VK region to the second conserved cysteine residue at position 88, adjacent to CDR III of these clones, namely CPVK46-1, CPVK46-3, CPVK46-4, CPVK46-5, CPVK46-6, CPVK46-7, CPVK46-8, CPVK46-11 and CPVK46-14 are shown in SEQ ID NOs: 28, 29, 30, 31, 32, 33, 34, 35 and 36, respectively. The amino acid sequences of the region encoding framework IV of these clones for CPVK46-6 and CPVK46-7 are shown in SEQ ID NOs: 86 and 87,

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The chimpanzee VK amino acid sequences comprising the mature N-terminus and the second conserved cysteine residue at position 88 were used as query sequences in computer homology searching of the Kabat database. The results of this analysis are shown in Table 2. In each case the closest match was with a human VK region, displaying between 68% (46-4/HKL310) and 97% (46-11/HKL106) sequence identity at the amino acid level. It is evident that the chimpanzee VK sequences are distinct from the collection of human VK found in the Kabat database.

The human subgroup homology is presented in Table 2. Of the four major human VK subgroups, matches were found for the two most frequently isolated, indicating that the chimpanzee VK repertoire is at least homologous to members of the majority of the human VK repertoire. Further sampling of the chimpanzee VK cDNA library will likely identify a greater diversity of chimpanzee VK regions, including ones homologous to the remaining two human VK subgroups (VKII and VKIV).

10			<b>Table 2</b> Overall Amino	
	Clone	Closest Match	Acid Homology	VH Subgroup Match
	46-1	HKL10C	85%	I.
	46-3	HKL 100	91	I
	46-5	HKL 100	91	I
15	46-7	HKL 100	81	I
	46-8	HKL 10N	90	I
	46-11	HKL 106	97	I
	46-14	HKL 100	92	I
	46-4	HKL 310	68	III
20	46-6	HKL 310	96	III

### Example 3

## Random cDNA Cloning and Sequence Analysis of Cynomolgus VH Regions

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Splenic RNA was recovered from a single donor cynomolgus monkey (Macaca cynomolgus) by means of standard laboratory practice. Cynomolgus heavy chain V regions were cloned from the total RNA using Marathon RACE methodology (Clontech, Palo Alto, CA, USA) following exactly the manufacturer's protocol using 3' Cg1 gene specific primers. After RACE PCR amplification, DNA bands of the expected size were excised from agarose gels, the DNA was purified and cloned into a plasmid vector. Although this cDNA library contains many distinct heavy V region clones, eight were selected randomly for sequence analysis. Complete nucleic acid sequences and predicted protein sequences of the Cynomolgus VH cDNA clones 2-1, 2-3, 2-4, 2-5, 2-6, 2-7, 2-8 and 2-10 are shown in SEQ ID NOs: 37, 38, 39, 40, 41, 42, 43 and 44, respectively. amino acid sequences of the region from the first amino acid of the mature VH region to the second conserved cysteine residue at position 92, adjacent to CDR III of these clones, namely CyVH2-1, CyVH2-3, CyVH2-4, CyVH2-5, CyVH2-6, CyVH2-7, CyVH2-8 and CyVH2-10 are shown in SEQ ID NOs: 45, 46, 47, 48,

49, 50, 51 and 52, respectively. The amino acid sequences of the region encoding framework IV of these clones for CyVH2-1, CyVH2-3, CyVH2-4, CyVH2-6, CyVH2-10 and CyVH2-5 are shown in SEQ ID NOs: 88, 89, 90, 91, 92 and 93, respectively.

The cynomolgus VH amino acid sequences from the mature N-terminus and the second conserved cysteine residue at position 92, adjacent to CDRIII, were used as query sequences in computer homology searching of the Kabat database. The results of this analysis are shown in Table 3. In each case the closest match was with a human VH region, displaying between 62% (2-6/ HHC20E) and 84% (2-5/ HHC20F) sequence identity at the amino acid level. It is evident that the cynomolgus VH sequences are distinct from the collection of human VH found in the Kabat database. Matches were found for each of the three major human VH subgroups, indicating that the cynomolgus VH repertoire includes at least some members homologous to each of the major human subgroups. The human subgroup homology is presented in Table 3.

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20			<b>Table 3</b> Overall Amino	
	Clone	Closest Match	Acid Homology	VH Subgroup Match
	2-4	HHC10Y	83%	I
	2-10	HHC20G	83	II
25	2-8	HHC20F	74	II
	2-6	HHC20E	62	II
	2-5	HHC20F	84	II
	2-3	HHC20F	75	II
	2-1	HHC316	71	III
30	2-7	HHC31C	81	III

The results show that the overall sequence identity between the cynomolgus and human VH regions ranged between 62 and 84% with a mean identity of 77%. Based on this observation, further sampling of the cynomolgus random VH library will likely provide a substantially greater diversity of VH sequences from which to choose optimum acceptor frameworks for each particular donor VH region.

## 40 Example 4 Random cDNA Cloning and Sequence Analysis of Cynomolgus V K

Cynomolgus light chain VK regions were cloned from the total splenic RNA using Marathon RACE methodology (Clontech,

Regions

Palo Alto, CA, USA) following exactly the manufacturer's protocol and CK 3' gene specific primers. After RACE PCR amplification, DNA bands of the expected size were excised from agarose gels, the DNA was purified and cloned into a plasmid vector. Although this cDNA library contains many distinct light chain Vk region clones, six were selected randomly for sequence analysis. Complete nucleic acid sequences and predicted protein sequences of the Cynomolgus VK cDNA clones 4-2, 4-3, 4-5, 4-6, 4-10 and 4-11 are shown in SEO ID NOs: 53, 54, 55, 56, 57 and 58, respectively. amino acid sequences of the region from the first amino acid of the mature VK region to the second conserved cysteine residue at position 88, adjacent to CDRIII, of these clones, namely  $CyV\kappa4-2$ ,  $CyV\kappa4-3$ ,  $CyV\kappa4-5$ ,  $CyV\kappa4-6$ ,  $CyV\kappa4-10$  and  $CyV\kappa4-$ 11 are shown in SEQ ID NOs: 59, 60, 61, 62, 63 and 64, respectively. The amino acid sequences encoding the framework IV region of these clones for CyVK4-3, CyVK4-6 and CyVk4-11 are shown in SEQ ID NOs: 94, 95 and 96, respectively.

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The cynomolgus VK amino acid sequences comprising the mature N-terminus and the second conserved cysteine residue at position 88 were used as query sequences in computer homology searching of the Kabat database. The results of this analysis are shown in Table 4. In each case the closest match was with a human Vk region, displaying between 73% (4-11/ HKL10S) and 94% (4-3/ HKL400) sequence identity at the amino acid level. It is evident that the cynomolgus VK sequences are distinct from the collection of human  $V\kappa$  found in the public genetic databases. The human subgroup homology is presented in Table 4. Matches were found for three of the four major human Vk subgroups, indicating that the cynomolgus VK repertoire is largely homologous to members of the majority of the human Vk repertoire. Further sampling of the cynomolgus Vk cDNA library will likely identify a greater diversity of cynomolgus VK regions, including ones homologous to the remaining human  $V\kappa$  subgroup ( $V\kappa III$ ).

Table 4
Overall Amino

	Clone	Closest Match	Acid Homology	Vκ Śubgroup Match
5	4-6	HKL10L	80%	I
	4-2	HKL10Z	83	I
	4-11	HKL10S	73	I
	4-10	HKL10F	93	I
	4-5	HKL209	86	II
10	4-3	HKL400	94	IV

The results show that the overall sequence identity between the cynomolgus and human VK regions ranged between 73 and 94% with a mean identity of 85%. Based on this observation, further sampling of the cynomolgus random VK library will provide a substantially greater diversity of VK sequences from which to choose optimum acceptor frameworks for each particular donor VK region.

20 Example 5

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### Preparation of Engineered Anti-IL-5 Monoclonal Antibodies

The VK and VH genes of the rat anti-interleukin-5 (IL-5) antibody 4A6 are shown in SEQ ID NOs: 65 and 66, respectively. These genes encode a high affinity neutralizing monoclonal antibody specific for human IL-5 useful for the treatment of asthma. See U.S. Patent No. 5,693,323.

The 4A6 light chain was engineered as follows. The sequence of donor antibody VK4A6 (SEQ ID NO: 65) was aligned with the acceptor antibody light chain VK region from the chimpanzee Mab C108G (Mol. Immunol. 32:1081-1092 (1995)) (SEQ ID NO: 67) as shown in Fig. 1. Since native VK4A6 has a unique deletion of residue 10, the sequence alignment included the insertion of a gap at that position. The CDR residues were identified as defined by the convention of Kabat et al. in Sequences of Proteins of Immunological Interest, 4th ed., U.S. Department of Health and Human Services, National Institutes of Health (1987).

Framework residues that could influence CDR presentation were identified by analysis of three-dimensional models based on known antibody crystal structures. The residues of this

CDR-contacting set were compared among the aligned VK4A6 and VKC108G sequences, and the positions of the set that differed between the VK4A6 and the VKC108G were marked (Fig. 1, asterisks). The CDRs and the marked framework residues of VK4A6 (the donor antibody) were transferred replacing the corresponding residues of VKC108G (the acceptor antibody). The completed engineered 4A6 light chain V region is shown in SEQ ID NO: 68. Six donor framework residues were retained in the engineered molecule at residues 1 to 4, 49 and 60.

In analogous fashion, a similar method was used to engineer the 4A6 heavy chain. The sequence of donor antibody VH4A6 (SEQ ID NO: 66) was aligned with the acceptor antibody heavy chain V region from the chimpanzee Mab C108G (SEQ ID NO: 69) as shown in Fig. 2. A large gap was introduced in the VH4A6 CDRIII alignment, as CDRIII of VHC108G is 10 residues longer. CDR residues were identified as defined by the convention of Kabat et al., supra.

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Framework residues that could influence CDR presentation were identified by analysis of three-dimensional models based on known antibody crystal structures. The residues of this CDR-contacting set were compared among the aligned VH4A6 and VHC108G sequences, and the positions of the set that differed between the VH4A6 and the VHC108G were marked (Fig. 2, asterisks). In total, 11 such CDR contacting residues that differed between VH4A6 and the VHC108G were selected and marked. The CDRs and the marked CDR contacting framework residues of VH4A6 (the donor antibody) were transferred replacing the corresponding residues of VHC108G (the acceptor The completed engineered 4A6 heavy chain V region antibody). is shown in SEQ ID NO: 70. Eleven donor framework residues were retained in the engineered molecule at residues 27, 30, 38, 49, 66, 67, 69, 71, 73, 78 and 94.

The engineered 4A6 can be expressed in cells using methods well known to those skilled in the art. Briefly, genes encoding the complete engineered 4A6 VH and VK regions can be assembled from long synthetic oligonucleotides and ligated into appropriate eukaryotic expression vectors containing the desired antibody constant regions. Such an expression vector will contain selectable markers, for

example, neomycin resistance and regulatory sequences, for example, the CMV promoter, required to direct the expression of full-length antibody heavy and light chains. Subsequently, transfection of the appropriate host cell, for example, chinese hamster ovary, would result in the expression of fully active engineered 4A6.

### Example 6

### Preparation of Engineered Anti-Integrin Monoclonal Antibodies

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The Vk and VH genes of the murine anti-integrin antibody B9 are shown in SEQ ID NOs: 71 and 72, respectively. These genes encode a high affinity neutralizing monoclonal antibody specific for human integrin  $\alpha v\beta 3$  useful for the treatment of vascular diseases.

The B9 light chain was engineered as follows. The amino acid sequence of donor antibody VkB9 (SEQ ID NO: 72) was compared to each of the nine chimpanzee Vk sequences described above and percent sequence identity determined by computer homology searching using the LASERGENE program "MEGALIGN" (DNASTAR, Inc., Madison, WI). Clones CPVk46-3 (SEQ ID NO: 29) and CPVk46-14 (SEQ ID NO: 36) were identified as the chimpanzee Vk regions with the highest overall sequence similarity (77%) to the B9 donor Vk. CPVk46-3 was selected as the acceptor framework.

Similarly, the chimpanzee Jk gene segment of CPVk46-1 (SEQ ID NO: 97) was selected as acceptor framework IV. The sequences of the donor VkB9 and acceptor CPVk46-3, CPVk46-1 V regions were aligned and the positions of their respective framework and CDRs were determined as shown in Fig. 3.

The CDR residues were identified as defined by the convention of Kabat *et al.*, *supra*. The results show that VKB9 and CPVK46-3 share 77% overall sequence identity, with the framework regions I through III sharing 81% sequence identity.

Framework residues that could influence CDR presentation were identified by analysis of three-dimensional models based on known antibody crystal structures. The residues of this

CDR-contacting set were compared among the aligned VkB9 and CPVk46-3 sequences, and none of this set were found that differed between the VkB9 and the CPVk46-3. Accordingly, only the CDRs of VkB9 (the donor antibody) were transferred replacing the corresponding residues of CPVk46-3 (the acceptor antibody). Lastly, the framework IV sequences of CPVk46-1 replaced the corresponding framework IV residues of the B9 light chain variable region. The completed engineered B9 light chain V region is shown in SEQ ID NO: 73. No donor framework residues were retained in the engineered light chain variable region.

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The B9 heavy chain was engineered in analogous fashion. The amino acid sequence of donor antibody VHB9 (SEQ ID NO: 71) was compared to each of the nine chimpanzee VH sequences described above by computer homology searching. Clone CPVH41-18 (SEQ ID NO: 17) was identified as the chimpanzee VH region with the highest overall sequence similarity (58%) to the B9 donor VH.

The chimpanzee JH gene segment of CPVH41-10 (SEQ ID NO: 82) was selected as acceptor framework IV. The sequences of the donor VHB9 and chimpanzee acceptor V regions were aligned and the positions of their respective framework and CDRs determined as shown in Fig. 4.

The CDR residues were identified as defined by the convention of Kabat et al., supra. The results show that VHB9 and CPVH41-18 share 58% overall sequence identity, with the framework regions I through III sharing 65% sequence identity.

Framework residues that could influence CDR presentation were identified by analysis of three-dimensional models based on known antibody crystal structures. The residues of this CDR-contacting set were compared among the aligned VHB9 and CPVH41-18 sequences, and the nine residues of the set that differed between VHB9 and the chimpanzee acceptor frameworks were marked. The CDRs and the marked framework residues of donor antibody VHB9 were transferred replacing the corresponding residues of CPVH41-18 (the acceptor antibody). Lastly, the framework IV sequences of CPVH41-10 replaced the corresponding framework IV residues of the B9 heavy chain variable region. The completed engineered B9 heavy chain

region is shown in SEQ ID NO: 74. Nine donor framework residues were retained in the engineered heavy chain variable region at positions 24, 27, 38, 48, 66, 67, 69, 93 and 94.

### Example 7

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## Expression and Characterization of Engineered Anti-Integrin Monoclonal Antibodies

The engineered B9 antibody was expressed in cells using methods well known to those skilled in the art. Briefly, genes encoding the complete engineered B9 VH and VK regions were assembled from long synthetic oligonucleotides and ligated into appropriate eukaryotic expression vectors containing IgG1,K antibody constant regions. The expression vector contained a selectable marker for neomycin resistance and CMV promoter regulatory sequences. Subsequent transfection of a COS host cell resulted in the expression of engineered B9 (CPB9).

The relative binding avidity of CPB9 was compared to that of the original murine B9 antibody as follows. antibodies present in culture supernatants from cells maintained in culture for 5 days after transfection with the expression constructs were compared to the parental murine B9 antibody using the ORIGEN technology (IGEN Inc, Gaithersburg, MD). Briefly, different dilutions of the B9 variants were incubated with purified human  $\alpha \nu \beta 3$  integrin which had previously been biotinylated, and an electrochemiluminescent TAG moiety specific for the antibody C regions. B9 variant antibody bound to the integrin was measured by capturing the immune complexes onto streptavidin beads followed by analysis on the ORIGEN instrument. The results showed that the CPB9 and the murine B9 binding curves were displaced only by about 3-fold indicating that the overall specific binding avidity of CPB9 and murine B9 for  $\alpha v \beta 3$  are within three-fold of each other. Accordingly, the results show that the CDR grafting of rodent CDRs onto chimpanzee frameworks as described in the present invention retained nearly all of the binding avidity of the parent rodent mAb.

### Example 8

## Preparation of Engineered Anti-Erythropoietin Receptor Monoclonal Antibodies

The VH and VK genes of the murine anti-erythropoietin receptor antibody 3G9 are shown in SEQ ID NOs: 75 and 76, respectively. These genes encode a high affinity neutralizing monoclonal antibody specific for human erythropoietin receptor (EPOr) useful for the treatment of hematopoietic disorders.

The 3G9 light chain was engineered as follows. The amino acid sequence of donor antibody Vκ3G9 (SEQ ID NO: 76) was compared to each of the nine chimpanzee Vκ sequences described above by computer homology searching as described above. Clones CPVκ46-3 (SEQ ID NO: 29), CPVκ46-5 (SEQ ID NO:

15 31), CPVκ46-8 (SEQ ID NO: 34) and CPVκ46-14 (SEQ ID NO: 36) were identified as the chimpanzee Vκ regions with the highest overall sequence similarity (65%) to the 3G9 donor Vκ.

CPVκ46-14 was selected as the acceptor framework.

The chimpanzee Jk gene segment of CPVk46-14 was

identical to that of CPVk46-1 (SEQ ID NO: 97) and was
selected as acceptor framework IV. The sequences of the
donor Vk3G9 and acceptor CPVk46-14 V regions were aligned and
the positions of their respective framework and CDRs were
determined as shown in Fig. 5.

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The CDR residues were identified as defined by the convention of Kabat et al., supra. The results show that VK3G9 and CPVK46-14 share 65% overall sequence identity, with the framework regions I through III sharing 73% sequence identity.

Framework residues that could influence CDR presentation were identified by analysis of three-dimensional models based on known antibody crystal structures. The residues of this CDR-contacting set were compared among the aligned VK3G9 and CPVK46-14 sequences, and the positions of this set that differed between VK3G9 and the CPVK46-3 were marked. The CDRs and marked residues of VK3G9 (the donor antibody) were

transferred replacing the corresponding residues of CPVK46-14 (the acceptor antibody). Lastly, the framework IV sequences of CPVK46-14 replaced the corresponding framework IV residues of the 3G9 light chain variable region. The completed engineered 3G9 light chain V region is shown in SEQ ID NO: 77. Three donor framework residues were retained in the engineered light chain variable region at positions 3, 46 and 60.

The 3G9 heavy chain was engineered in analogous fashion.

The amino acid sequence of donor antibody VH3G9 (SEQ ID NO:

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The chimpanzee JH gene segment of CPVH41-18 was identical to CPVH41-9 (SEQ ID NO: 81) and was selected as acceptor framework IV. The sequences of the donor VH3G9 and chimpanzee acceptor V regions were aligned and the positions of their respective framework and CDRs determined as shown in Fig. 6.

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The CDR residues were identified as defined by the convention of Kabat et al., supra. The results show that VH3G9 and CPVH41-18 share 53% overall sequence identity, with the framework regions I through III sharing 62% sequence identity.

Framework residues that could influence CDR presentation were identified by analysis of three-dimensional models based on known antibody crystal structures. The residues of this CDR-contacting set were compared among the aligned VH3G9 and CPVH41-18 sequences, and the twelve residues of the set that differed between VH3G9 and the chimpanzee acceptor frameworks were marked. The CDRs and the marked framework residues of donor antibody VH3G9 were transferred replacing the corresponding residues of CPVH41-18 (the acceptor antibody). Lastly, the framework IV sequences of CPVH41-18 replaced the corresponding framework IV residues of the 3G9 heavy chain variable region. The completed engineered 3G9 heavy chain V region is shown in SEQ ID NO: 78. Twelve donor framework residues were retained in the engineered heavy chain variable

region at positions 24, 27, 30, 38, 48, 66-69, 71, 73, and 94.

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### Example 9

### Expression and Characterization of Engineered anti-Erythropoietin Receptor Monoclonal Antibodies

The engineered 3G9 antibody was expressed in cells using methods well known to those skilled in the art. Briefly, genes encoding the complete engineered 3G9 VH and VK regions were assembled from long synthetic oligonucleotides and ligated into appropriate eukaryotic expression vectors containing IgG1,  $\kappa$  antibody constant regions. The expression vector contained a selectable marker for neomycin resistance and CMV promoter regulatory sequences. Subsequent transfection of COS host cells resulted in the expression of engineered 3G9 (CP3G9).

Culture supernatants from COS cells transiently transfected with chimpanzee framework engineered 3G9 were compared with another 3G9 variant for the ability to bind human EPOr. The entire extracellular domain of the EPOr was expressed as recombinant protein, purified, and adsorbed onto the wells of ELISA plates. Dilutions of different antibodies were then tested for the ability to specifically bind to the solid phase associated EPOr.

HZ3G9 is a humanized variant of 3G9 in which human frameworks were used in traditional CDR grafting experiments. The humanized 3G9 heavy chain amino acid sequence is shown in SEQ ID NO: 79. The humanized 3G9 light chain sequence is shown in SEQ ID NO: 80. Previous experiments showed that HZ3G9 retained the full binding affinity and avidity of the parental murine 3G9. Accordingly, since HZ3G9G1 is identical to the chimpanzee version in all respects except the V region cassette, it was used in the present comparative binding experiments as a surrogate for murine 3G9. Negative control antibodies were also tested, including HZD12 which is a humanized antibody specific for human integrin, and CPB9 which is a chimpanzee framework engineered antibody specific for human integrins described above. Different concentrations of the 3G9 variants and control antibodies were incubated for one hour. After washing, the bound

antibodies were detected by incubation with anti-human H+L antibody-enzyme conjugate, a final wash, and addition of chromagen.

The binding curves obtained for CP3G9 and HZ3G9 were superimposable. This result indicates that the human and the chimpanzee framework engineered versions of 3G9 have identical overall binding avidity for the specific antigen human EPOr. Since the constant regions of HZ3G9 and CP3G9 are identical, the results also suggest the full binding affinity of the original rodent 3G9 is retained in the chimpanzee version of 3G9. Accordingly, the results show that CDR grafting of rodent CDRs onto chimpanzee acceptor frameworks as described in the present invention retained the full binding avidity of the parental rodent antibody.

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A BIAcore analysis (Pharmacia) was performed to determine the binding affinity for human EPOr of murine 3G9 and CP3G9. The interaction of CP3G9 as well as murine 3G9 with EPOr was characterized using a BIAcore 1000 biosensor. Descriptions of the instrumentation and the sensor surfaces are described in Brigham-Burke et al., Anal. Biochem., 205:125-131 (1992).

CP3G9 was captured onto a sensor surface of immobilized protein A. The kinetic binding constants were determined by passing solutions of monomeric EPOr over the surface and monitoring binding versus time. The equilibrium dissociation constant for the interaction was then derived from the ratio of the kinetic constants. The parent murine 3G9 was captured onto a surface of protein A captured rabbit anti-mouse Fc specific polyclonal antibody. The kinetics and dissociation constant for the interaction with EPOr was determined as described above. All measurements were made in 10 mM sodium phosphate, 150 mM NaCl pH 7.2 3 mM EDTA and 0.005% Tween 20. The flow rate was 60 uL/min. The temperature was 20° C.

	$k_{ass}$ $(M^{-1}s^{-1})$	kdiss (s-1)	K <sub>D</sub> (nM)
murine 3G9	1.2x106	$4.0 \times 10^{-3}$	3.3
CP3G9	$1.0 \times 10^{6}$	9.1x10 <sup>-3</sup>	9.1

These results show that the dissociation equilibrium constants determined for the murine and chimpanzee framework versions of 3G9 are within three fold of each other. This

data is in good agreement with the results of the ELISA-based study described above. Accordingly, the results show that the process used in generating the chimpanzee version of 3G9 largely retained the binding affinity of the original rodent mAb.

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The present invention may be embodied in other specific forms without departing from the spirit or essential attributes thereof, and, accordingly, reference should be made to the appended claims, rather than to the foregoing specification, as indicating the scope of the invention.

### Claims

- 1. An antibody comprising donor CDRs derived from an antigen-specific donor antibody of a non-human species and acceptor framework residues derived from a non-human primate.
- 2. The antibody of claim 1 wherein the non-human primate is an Old World ape.
- 3. The antibody of claim 2 wherein the Old World ape is Pan troglodytes, Pan paniscus or Gorilla gorilla.
- 4. The antibody of claim 3 wherein the Old World ape is Pan troglodytes.
- 5. The antibody of claim 1 further comprising one or more CDR-contacting residues of the donor antibody.
- 6. The antibody of claim 1 comprising human or Old World ape constant regions.
- 7. The antibody of claim 1 wherein one or more solvent-exposed framework residues are replaced with corresponding residues from a homologous selected non-human primate framework.
- 8. The antibody of claim 1 wherein the non-human primate is an Old World monkey.
- 9. The antibody of claim 8 wherein the Old World monkey genus is Macaca.
- 10. The antibody of claim 9 wherein the Old World monkey is Macaca cynomolgus.
- 11. The antibody of claim 8 further comprising one or more CDR-contacting residues of the donor antibody.
- 12. The antibody of claim 8 comprising human or Old World ape constant regions.

13. The antibody of claim 8 wherein one or more solvent-exposed framework residues are replaced with corresponding residues from a homologous selected non-human primate framework.

- 14. A method for making an antibody having reduced immunogenicity in humans comprising grafting CDRs from antigen-specific non-human antibodies onto homologous Old World ape acceptor frameworks.
- 15. The method of claim 14 wherein the Old World apeacceptor framework is from Pan troglodytes, Pan paniscus or Gorilla gorilla.
- 16. The method of claim 15 wherein the Old World ape acceptor framework is from Pan troglodytes.
- 17. A method for making an antibody having reduced immunogenicity in humans comprising grafting CDRs from antigen-specific non-human antibodies onto homologous Old World monkey acceptor frameworks.
- 18. The method of claim 17 wherein the Old World monkey acceptor framework is from the genus Macaca.
- 19. The method of claim 18 whereiin the Old World Monkey acceptor framework is from Macaca cynomolgus.
- 20. A chimpanzee VH acceptor framework I, II and III comprising an amino acid sequence as set forth in SEQ ID NOs: 10, 11, 12, 13, 14, 15, 16, 17 or 18.
- 21. A chimpanzee VH acceptor framework IV comprising an amino acid sequence as set forth in SEQ ID NOs: 81, 82, 83, 84 or 85.
- 22. A chimpanzee  $V\kappa$  acceptor framework I, II and III comprising an amino acid sequence as set forth in SEQ ID NOs: 28, 29, 30, 31, 32, 33, 34, 35 or 36.

23. A chimpanzee VK acceptor framework IV comprising an amino acid sequence as set forth in SEQ ID NOs: 86 or 87.

- 24. A cynomolgus VH acceptor framework I, II and III comprising an amino acid sequence as set forth in SEQ ID NOs: 45, 46, 47, 48, 49, 50, 51 or 52.
- 25. A cynomolgus VH acceptor framework IV comprising an amino acid sequence as set forth in SEQ ID NOs: 88, 89, 90, 91, 92 or 93.
- 26. A cynomolgus VK acceptor framework I, II and III comprising an amino acid sequence as set forth in SEQ ID NOs: 59, 60, 61, 62, 63 or 64.
- 27. A cynomolgus VK acceptor framework IV comprising an amino acid sequence as set forth in SEQ ID NOs: 94, 95 or 96.
- 28. An isolated nucleic acid molecule encoding the amino acid sequence of SEQ ID NOs: 10, 11, 12, 13, 14, 15, 16, 17, 18, 28, 29, 30, 31, 32, 33, 34, 35 or 36.
- 29. An isolated nucleic acid molecule encoding the amino acid sequence of SEQ ID NOs: 81, 82, 83, 84, 85, 86 or 87.
- 30. An isolated nucleic acid molecule encoding the amino acid sequence of SEQ ID NOs: 45, 46, 47, 48, 49, 50, 51, 52, 59, 60, 61, 62, 63 or 64.
- 31. An isolated nucleic acid molecule encoding the amino acid sequence of SEQ ID NOs: 88, 89, 90, 91, 92, 93, 94, 95 or 96.

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### Figure 1

4A6 DTVLTQSPA. LAVPPGERVT VSC**RASESVS TFLH**WYQQKP GHQP C108G AVHMTQSPSS LSASVGDSVT ITC**RASQTIN IYLN**WYQQKP GKAP

4A6 KLLIY**LASKL ES**GVPARFSG GGSGTDFTLT IDPVEADDTA TYYC**QQTWND** C108G KLLIF**DASIL QS**GVPSRFSG SGSGTDFSLT IRSLQPEDFA TYYC**QCGWGTH** 

4A6 PRTFGGGT KLELKR C108G PYNFGQGT KLEIKR 2 / 6

### Figure 2

4A6 EVQLQQSGPE VGRPGSSVKI SCKASGYTFT **DYVLM**VK QSPGQGLEWI C108G EVQLVESGGG VVQPGGSLRL SCAASGFTFD **DFAMH**WVR QAPGKGLEWI

4A6 GWIDPDYG TTDYAEKFKK KATLTADTSS STAYIQLSSL TSEDTATYFC C108G SLVSWDSY NIYHADSVKG RFTISRDNSR NSLYLQMNDL RPEDTAIYFC

4A6 AR**SRNYGG.....YI MY**WGQGVMVTVS C108G AK**ADTGGDFD YVSDSWRCAL DY**WGQGTLVTVS 3 / 6

### Figure 3

	1		C	DR1	
VLB9	DIQMTQTTSS	LSASLGDRVT	ITCRSSQ	DISNFLN	WYQQKPDGTV
Cmp46-3	DIQMTQSPSS	LSASVGDRVT	ITCRASQ	GISNYLA	WYQQKPGKAP
	45 <b>CDR2</b>				<b>CDR3</b> 94
VLB9	KLLIY <b>YTSTL</b>	<i>HS</i> GVPSRFSG	SGSGTDYSLT	ISNLEQEDIA	TYFC QQGNTL
Cmp46-3	KLLIY <b>YASRL</b>	<i>ES</i> GVPSRFSG	SGSGTDYTLT	ISSLQPEDFA	TYYC <i>qq<b>ynsn</b></i>
	95				
VLB9	PWTFGGGT	NLEIKR			
cmp46-1	FGGGT	KVEIKR			

4 / 6

Figure 4

CDR1 48 11 QVQLQQSGAE LMKPGASVKI SCKATGYTFS SYWIE..WVK QRPGHGLEWI VHB9 AMP41CL18 QVQLVQSGAE VKKPGSSVKV SCKVSGGTFS TYGFS..WVR QAPGQGLEWM 76 83 92 49 CDR2 66 . \*\* \* GEILP. RSG NTNYNEKFKG KATFTAETSS NTAYMQLSSL TPEDSAVYYC VHB9 AMP41CL18 GMIIP..IVG TVKYAQRFQG RVSINADTST NIAYMELTSL RSEDTAVYYC 93 CDR3 104 SSRGVRGSM.....DYW GQGTSVTVSS AMP41CL18 ATDLTVTTNDAF....DI AMP41CL10 W GOGTLVTVSS

5 / 6

## Figure 5

	1		C	DR1	
VL3G9 VK46-14		MSTSVGDRVS LSASVGDRVT			
	45 <i>CDR2</i>	*			<b>CDR3</b> 94
VL3G9 VK46-14	******	<i>YS</i> GVPDRFTG <i>QS</i> GVPSRFSG		ISNVQSEDLA ISSLQPEDFA	~
VL3G9 VK46-14	95 <b>plt</b> fgagt <b>hpt</b> fgggt				

6 / 6

## Figure 6

	1	11	21	CDR1	39 48
VH3G9 Chimp41-18			SCKASGSTFT SCKVSGGTFS		QRPGRGLEWI
4	19	CDR2	66 7	76	83 92
VH3G9 Chimp41-18			KATLTVDKPS RVSINADTST		
9	3 <b>CDR3</b>	:	L0 <b>4</b>		
VH3G9 Chimp41-18	AR <b>ETYYDSS.</b> AT <b>DLTVTTN.</b>	FAYW	~		

## SEOUENCE LISTING

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Pro Ser Gln Thr Leu Ser Leu Thr Cys Ala Val Ser Gly Gly Ser Ile

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tac tac ttt gac tac tgg ggc ccg gga acc ctg gtc acc gtc ttc

429

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_	_		ttc						_		_					220
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110	- 7 -	115	Cys	ALG	птэ	nr 9	120		DCI	501	пор	125	1110	2135	1 110	
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105

100

110

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ggt	gcc	aga	tgt	gac	atc	cag	atg	acc	cag	ttt	cca	tcc	tcc	ctg	tct	96
Gly	Ala	Arg	Cys	Asp	Ile	Gln	Met	Thr	Gln	Phe	Pro	Ser	Ser	Leu	Ser	
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gca	tct	gta	gga	gac	aga	gtc	acc	atc	act	tgc	cag	tca	agt	cag	agc	144
Ala	Ser	Val	Gly	Asp	Arg	Val	Thr	Ile	Thr	Cys	Gln	Ser	Ser	Gln	Ser	
		35					40					45				
						tgg										192
Ile		Asn	Cys	Leu	Ser	Trp	Tyr	Gln	Gln	Lys		Gly	Lys	Ala	Pro	
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	a+ a	a+ -	5 t a	<del>+-+</del>	~~+	gca	ttc	=00	++~	aat	agt	aaa	atc	cca	tca	240
						Ala										240
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Arg	Phe	Ser	Gly	Ser	Gly	Ser	Gly	Thr	Asp	Phe	Thr	Leu	Thr	Ile	Ser	
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						ttt										336
Asn	Leu	Gln	Pro	Glu	Asp	Phe	Ala	Thr	Tyr	Tyr	Сув	Gln		Gly	Tyr	
			100					105					110			
																201
						ggt										381
GIĀ	Thr	115	ьeu	Thr	Pne	Gly	120	GIY	THE	пур	vaı	125	116	nys		
		113					120									
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ctc	cca	ggt	acc	aga	tgt	gac	atc	cag	atg	acc	cag	tct	cca	tcc	tcc	96
Leu	Pro	Gly	Thr	Arg	Cys	Asp	Ile	Gln	Met	Thir	Gln	Ser	Pro	Ser	Ser	
			20					25					30			
ctg	tct	gca	tct	gta	gga	gac	aga	gtc	acc	atc	act	tgc	cgg	gcc	agt	144
Leu	Ser	Ala	Ser	Va1	Gly	Asp	Arg	Va1	Thr	Ile	Thr	Cys	Arg	Ala	Ser	
		35					40					45				
cag	ggc	att	agc	aat	tat	tta	gcc	tgg	tat	cag	cag	aaa	cca	ggg	aaa	192
Gln	Gly	Ile	Ser	Asn	Tyr	Leu	Ala	Trp	Tyr	Gln	Gln	Lys	Pro	Gly	Lys	
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gcc	cct	aag	ctc	ctc	atc	tat	tat	gca	tcc	aga	ttg	gaa	agt	ggg	gtc	240
Ala	Pro	Lys	Leu	Leu	Ile	туг	Tyr	Ala	Ser	Arg	Leu	Glu	Ser	Gly	Val	
65					70					75					80	
cca	tca	agg	ttc	agc	ggc	agt	gga	tct	ggg	acg	gat	tac	act	ctc	acc	288
Pro	Ser	Arg	Phe	Ser	Gly	Ser	Gly	Ser	Gly	Thr	Asp	Tyr	Thr	Leu	Thr	
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atc	agc	agc	ctg	cag	cct	gaa	gat	ttt	gca	act	tat	tac	tgt	caa	cag	336
Ile	Ser	Ser	Leu	Gln	Pro	Glu	Asp	Phe	Ala	Thr	Tyr	Tyr	Cys	Gln	Gln	
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tat	aac	agt	aac	ccc	ttt	tcg	gtg	gag	gga	cca	agg	tgg	aga	tca	aac	384
Tyr	Asn	Ser	Asn	Pro	Phe	Ser	Val	Glu	Gly	Pro	Arg	Trp	Arg	Ser	Asn	
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105

100

110

tta	cct	cat	acg	ctc	act	ttc	ggt	gga	ggg	acc	aag	gtg	gag	atc	aaa	384
Leu	Pro	His	Thr	Leu	Thr	Phe	Gly	Gly	Gly	Thr	Lys	Vail.	Glu	Ile	Lys	
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1				5					10					15		
												ctg				96
Arg	Cys	Asp		Gln	Met	Thr	Gln		Pro	Ser	Ser	Leu		Ala	Ser	
			20					25					30			
												cag				144
Val	GTĀ		Arg	Val	Thr	Ile		Cys	Gin	Ala	Ser	Gln	ser	TTE	Ser	
		35					40					45				
										~~~	227	~~~	aat	220	ata	192
												gcc				132
ASII		ьeu	ser	TID	TYL		GIII	ьуѕ	PIO	Сту	60 Dys	Ala	PIO	пур	nea	
	50					55					00					
a+~	2+-	+-+	~=+	~~-	+~~	act	++~	C22	act	מממ	ata	cca	tc=	acc	ttc	240
												cca Pro				240
	тте	TAL	ASP	wrg		1111	ьеи	<b>⇔</b> ±11	ner	75	val		≏e1	r.a	80	
65					70					, ,					30	
a.c.+	aaa	a c+	aa=	+ = +	aaa	202	ϱ	ttc	act	ctc	acc	atc	age	act	cta	288
agı	990	ayı	yya		999	aça	yac		با ب		ucc	سدد	ug-	uyı	c-g	200

Ser Gly Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile Ser Ser Leu 90 85 caa cct gaa gat ttt gca aca tat tac tgt cag cgt ggt tac ggt aca 336 Gln Pro Glu Asp Phe Ala Thr Tyr Tyr Cys Gln Arg Gly Tyr Gly Thr 105 110 100 372 ctc act ttc ggt gga ggg acc aag gtg gag atc aaa Leu Thr Phe Gly Gly Gly Thr Lys Val Glu Ile Lys 120 115 <210> 23 <211> 384 <212> DNA <213> Pan troglodytes <220> <221> CDS <222> (1)...(384) <400> 23 atg gaa gcc cca gcg cag ctt ctc ttc ctc ctg cta ctc tgg ctc cca 48 Met Glu Ala Pro Ala Gln Leu Leu Phe Leu Leu Leu Leu Trp Leu Pro 15 10 1 5 gat acc acc gga gaa ata gtg ttg acg cag tct cca gcc acc ctg tct Asp Thr Thr Gly Glu Ile Val Leu Thr Gln Ser Pro Ala Thr Leu Ser 30 20 ttg tct cca ggg gaa aga gcc acc ctc tcc tgc agg gcc agt cag agt 144 Leu Ser Pro Gly Glu Arg Ala Thr Leu Ser Cys Arg Ala Ser Gln Ser 40 35 gtt agc agg tac tta gcc tgg tac cag cag aaa cct ggc cag gct ccc 192 Val Ser Arg Tyr Leu Ala Trp Tyr Gln Gln Lys Pro Gly Gln Ala Pro 60 50 55

agg	ctc	ctc	atc	tat	ggt	gca	tcc	aac	agg	gcc	act	ggc	atc	cca	gcc	240
Arg	Leu	Leu	Ile	$\mathtt{Tyr}$	Gly	Ala	Ser	Asn	Arg	Ala	Thr	Gŀ.	-Ile	Pro	Ala	
65					70					75					80	
agg	ttc	agt	ggc	agt	ggg	tct	agg	aca	gac	ttc	act	ctc	acc	atc	agc	288
Arg	Phe	Ser	Gly	Ser	Gly	Ser	Arg	Thr	Asp	Phe	Thr	Leu	Thr	Ile	Ser	
				85					90					95		
													cag			336
Ser	Val	GLu		Glu	Asp	Phe	Ala		Tyr	ТУĽ	Cys	GIN	Gln	туr	Asn	
			100					105					110			
330	C20	aat	ata	ato	acc	++-	ממם	caa	aaa	aca	cas	cta	gag	att	222	384
													Glu			301
ASII	GIII	115	Leu	116	A.Lu	1110	120	O I I I	OLY	1111	******	125			230	
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Met 1	<2 <2 <2 <4 gac Asp	213> 220> 221> 222> 100> atg	CDS (1) 24 agg Arg	gtc Val	387) ccc Pro	gct Ala	Gln	Leu	Leu 10	Gly	Leu	Leu	Leu	Leu 15	Trp	
Met 1 ttc	<2 <2 <2 <2 gac Asp	213> 220> 221> 222> 100> atg Met	Pan CDS (1) 24 agg Arg	gtc Val 5	387) ccc Pro	gct Ala gac	Gln	Leu cag	Leu 10 atg	Gly acc	Leu cag	Leu tct	Leu	Leu 15 tcc	Trp	<b>4</b> 8
Met 1 ttc	<2 <2 <2 <2 gac Asp	213> 220> 221> 222> 100> atg Met	Pan CDS (1) 24 agg Arg	gtc Val 5	387) ccc Pro	gct Ala gac	Gln	Leu cag Gln	Leu 10 atg	Gly acc	Leu cag	Leu tct	cct Pro	Leu 15 tcc	Trp	
Met 1 ttc	<2 <2 <2 <2 gac Asp	213> 220> 221> 222> 100> atg Met	Pan CDS (1) 24 agg Arg	gtc Val 5	387) ccc Pro	gct Ala gac	Gln	Leu cag	Leu 10 atg	Gly acc	Leu cag	Leu tct	Leu	Leu 15 tcc	Trp	
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25

Leu Ser Ala Ser Ile Gly Asp Arg Val Thr Ile Thr Cys Arg Ala Ser 40 35 cag ggc atc tat aat tat ttg aat tgg tat cag caa aaa cca ggg aga 192 Gln Gly Ile Tyr Asn Tyr Leu Asn Trp Tyr Gln Gln Lys Pro Gly Arg 55 60 50 gcc cct gga ctc ctc atc ttt ggt gcc agg aat ttg gag act ggg gtc 240 Ala Pro Gly Leu Leu Ile Phe Gly Ala Arg Asn Leu Glu Thr Gly Val 75 80 70 65 cca tca aca ttc agc ggc agt ggt tcc ggg aca cac ttc act ctc acc 288 Pro Ser Thr Phe Ser Gly Ser Gly Ser Gly Thr His Phe Thr Leu Thr 90 95 85 atc agc agc ctg cag cct ggt gat ttt gcg act tat tac tgt cag caa 336 Ile Ser Ser Leu Gln Pro Gly Asp Phe Ala Thr Tyr Tyr Cys Gln Gln 110 105 100 tat tat act acc ccg tat act ttt ggc cag ggg acc aag ctg gag atc 384 Tyr Tyr Thr Thr Pro Tyr Thr Phe Gly Gln Gly Thr Lys Leu Glu Ile 125 120 115 387 aaa <210> 25 <211> 387 <212> DNA <213> Pan troglodytes <220> <221> CDS <222> (1)...(387) <400> 25 atg gac atg agg gtc ccc gct cag ctc ctg ggg ctc ctg ctg ctc tgt 48 Met Asp Met Arg Val Pro Ala Gln Leu Leu Gly Leu Leu Leu Cys

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Phe	Pro	Gly	Ala	Arg	Cys	Asp	Ile	Gln	Met	Thr	Gln	Ser	Pro	Ser	Ser	
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Leu	Ser	Ala	Ser	Va1	Gly	Asp	Arg	Va1	Thr	Ile	Ser	Cys	Arg	Ala	Ser	
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										act						240
	Pro	гÀЗ	Pro	Leu		ıyr	Ala	Ala	ser	Thr	reu	PFO	ser	GΙΆ		
65					70					75					80	
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Pro	Ser	Arg	Phe	Ser	Gly	Ser	Gly	Ser	Gly	Thr	Asp	Phe	Thr	Leu	Thr	
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atc	agc	agc	ctg	cag	cct	gaa	gat	tct	gca	act	tat	tac	tgc	cga	caa	336
Ile	Ser	Ser	Leu	Gln	Pro	Glu	Asp	Ser	Ala	Thr	Tyr	Tyr	Cys	Arg	Gln	
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tat	aat	agt	tat	ccg	ctc	act	ttc	ggt	gga	ggg	acc	aag	gtg	gag	atc	384
Tyr	Asn	Ser	Tyr	Pro	Leu	Thr	Phe	Gly	G1y	Gly	Thr	Lys	Val	Glu	Ile	
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372 cga acg ttc ggc caa ggg acc aag ctg gaa atc aaa Arg Thr Phe Gly Gln Gly Thr Lys Leu Glu Ile Lys 115 120

110

Pro Asp Asp Phe Ala Thr Tyr Tyr Cys Gln Gln Tyr Ser Ser Tyr Pro 105

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Ile Ser Ser Leu Gln Pro Glu Asp Phe Ala Thr Tyr Tyr Cys Gln His

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ggt tac ggt aca cat ecc act tte ggt gga ggg ace aag gtg gag ate 384
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aaa 387

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Leu Ser Trp Tyr Gln Gln Lys Pro Gly Lys Ala Pro Thr Leu Leu Ile 35 40 45

Tyr Gly Ala Phe Thr Leu Asn Ser Gly Val Pro Ser Arg Phe Ser Gly 50 55 60

Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile Ser Asn Leu Gln Pro 65 70 75 80

Glu Asp Phe Ala Thr Tyr Tyr Cys

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Glu Lys Val Thr Ile Thr Cys Arg Ala Ser Gln Ser Ile Gly Ser Ser
20 25 30

Leu His Trp Tyr Gln Gln Lys Pro Gly Gln Ser Pro Lys Leu Leu Ile 35 40 45

Lys Tyr Ala Ser Gln Ser Ile Ser Gly Val Pro Ser Arg Phe Ser Gly 50 55 60

Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile Asn Ser Leu Glu Ala 65 70 75 80

Glu Asp Ala Ala Thr Tyr Tyr Cys

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1 5 10 15

Asp Arg Val Thr Ile Thr Cys Gln Ala Ser Gln Ser Ile Ser Asn Tyr
20 25 30

32

Leu Ser Trp Tyr Gln Gln Lys Pro Gly Lys Ala Pro Lys Leu Leu Ile 40 Tyr Asp Ala Ser Thr Leu Gln Ser Gly Val Pro Ser Arg Phe Ser Gly 60 55 Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile Ser Ser Leu Gln Pro 65 70 75 80 Glu Asp Phe Ala Thr Tyr Tyr Cys 85 <210> 32 <211> 88 <212> PRT <213> Pan troglodytes <220> <221> DOMAIN <222> (24)...(34) <223> CDRI <221> DOMAIN <222> (50)...(66) <223> CDRII <400> 32 Glu Ile Val Leu Thr Gln Ser Pro Ala Thr Leu Ser Leu Ser Pro Gly 10 Glu Arg Ala Thr Leu Ser Cys Arg Ala Ser Gln Ser Val Ser Arg Tyr 25 30 20 Leu Ala Trp Tyr Gln Gln Lys Pro Gly Gln Ala Pro Arg Leu Leu Ile 40 35 Tyr Gly Ala Ser Asn Arg Ala Thr Gly Ile Pro Ala Arg Phe Ser Gly 60 55 Ser Gly Ser Arg Thr Asp Phe Thr Leu Thr Ile Ser Ser Val Glu Pro 80 75 65 70

85

Glu Asp Phe Ala Val Tyr Tyr Cys

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1 5 10 15

Asp Arg Val Thr Ile Ser Cys Arg Ala Ser Leu Asp Ile Ser Thr Trp
20 25 30

Leu Ala Trp Tyr Gln Gln Lys Pro Gly Lys Ala Pro Lys Pro Leu Ile 35 40 45

Tyr Ala Ala Ser Thr Leu Pro Ser Gly Val Pro Ser Arg Phe Ser Gly 50 55 60

Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile Ser Ser Leu Gln Pro 65 70 75 80

Glu Asp Ser Ala Thr Tyr Tyr Cys

85

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1 5 10 15

Asp Arg Val Thr Ile Thr Cys Arg Ala Ser Gln Gly Ile Ser Ser Trp

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Ser Leu Tyr Leu Glu Met Asn Ser Leu Arg Pro Asp Asp Thr Ala Val

100 105 110

tat ttc tgt gtg aga gaa tac aga gat gga ctg gat gtc tgg ggc cgg 384
Tyr Phe Cys Val Arg Glu Tyr Arg Asp Gly Leu Asp Val Trp Gly Arg

115 120 125

gga gtt ctg gtc acc gtc tcc tca 408
Gly Val Leu Val Thr Val Ser Ser

130 135

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ggc cca gga ctg gtg aag cct tcg gag acc ctg tcc ctc act tgt act

Gly Pro Gly Leu Val Lys Pro Ser Glu Thr Leu Ser Leu Thr Cys Thr

20

25

30

gtc tct ggt gac tcc atc acc act gtc ttc tgg agc tgg ctc cgc cag

144

Val Ser Gly Asp Ser Ile Thr Thr Val Phe Trp Ser Trp Leu Arg Gln

35

40

45

tcg cca ggg att ggg ctg gag tgg att ggg aat ttt gct ggt agt act

192

Ser Pro Gly Ile Gly Leu Glu Trp Ile Gly Asn Phe Ala Gly Ser Thr

50

55

60

35 40 45 acc gac agc tgg atc agc tgg gtg cgc cag atg ccc ggg aaa ggc ctg 192 Thr Asp Ser Trp Ile Ser Trp Val Arg Gln Met Pro Gly Lys Gly Leu 50 55 60 gag tgg atg gga aac atc tat cct ggt gat tct gat tcc aga tac aac 240 Glu Trp Met Gly Asn Ile Tyr Pro Gly Asp Ser Asp Ser Arg Tyr Asn 80 65 70 288 ccg tcc ttc caa ggc cgc gtc act atc tca gtc gac aag tcc atc agt Pro Ser Phe Gln Gly Arg Val Thr Ile Ser Val Asp Lys Ser Ile Ser 90 95 85 ace ace tac etg cag tgg age age etg aag gee teg gae act gee aca 336 Thr Thr Tyr Leu Gln Trp Ser Ser Leu Lys Ala Ser Asp Thr Ala Thr 105 110 100 tat tac tgt gcg aag ata gat agc aac tac tac agc cgg ttc gaa gtc 384 Tyr Tyr Cys Ala Lys Ile Asp Ser Asn Tyr Tyr Ser Arg Phe Glu Val 125 115 120 417 tgg ggc ccc gga gtc atg gtc acc gtc tcc tca Trp Gly Pro Gly Val Met Val Thr Val Ser Ser 130 135 <210> 40 <211> 423 <212> DNA <213> Macaca cynomolgus <220> <221> CDS

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	Lys															
	пуз	1115	пец	5	1110	1110	200	200	10		****			15		
1				5					10					1.0		
									****	~~~		~~~	~ + · ~	~ ~ ~	~	96
	ctg -															90
Val	Leu	Ser		Val	GIn	Leu	Gin		ser	GIA	Pro	GTĀ		val	ьуs	
			20					25					30			
	tcg															144
Pro	Ser	Glu	Thr	Leu	Ser	Leu	Thr	Cys	Thr	Val	Ser	Gly	Gly	Ser	Phe	
		35					40					45				
agt	act	tac	tac	tgg	aat	tgg	atc	cgc	cag	CCC	cca	aaa	aag	gga	ctg	192
Ser	Thr	Tyr	Tyr	Trp	Asn	Trp	Ile	Arg	Gln	Pro	Pro	Gly	Lys	Gly	Leu	
	50					55					60					
gag	tgg	att	gga	tat	atc	ggt	ggt	ggt	ggt	ggt	cgc	ccc	aac	tac	aat	240
Glu	Trp	Ile	Gly	Tyr	Ile	Gly	Gly	Gly	Gly	G1y	Arg	Pro	Asn	Tyr	Asn	
65					70					75					80	
tcc	tcc	ctc	aag	agt	cgc	atc	acc	ctg	tca	cta	gac	gcg	tcc	aag	aac	288
Ser	Ser	Leu	Lys	Ser	Arg	Ile	Thr	Leu	Ser	Leu	Asp	Ala	Ser	Lys	Asn	
				85					90					95		
caq	ttc	tcc	ctq	aac	ctg	agc	tct	gtg	acc	gcc	gcg	gaç	acg	gcc	gtg	336
_	Phe		_													
			100					105					110			
tac	tac	tat	acc	aga	gat	caa	aac	tac	aat	acc	agc	aat	gat	act	ttt	384
	Tyr															
-1-	-2-	115				3	120	-4-				125	-			
		<b>-</b>			<b>a</b>	a+ c	2~~	ata	200	at-a	tat	tes				423
	ttc															423
Asp	Phe	_	GīĀ	GIN	ĠΤĀ			val	THE	val		ser				
	130					135					140					

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tat	tac	tgt	gtc	aga	tcg	acg	gca	tta	ttt	tcg	ttg	gat	gtc	tgg	ggc	384
Tyr	Tyr	Cys	Val	Arg	Ser	Thr	Ala	Leu	Phe	Ser	Leu	Asp	Val	Trp	Gly	
		115					120					125				
cgg	gga	ctt	ctg	gtc	acc	gtc	tcc	tca								411
Arg	Gly	Leu	Leu	Val	Thr	Val	Ser	Ser								
	130					135										
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1				5					10					15		
	gtc															96
Gly	Val	Gln		Asp	Lys	Gln	Leu		Gin	Ser	GLY	GLY		Leu	Val	
			20					25		ė			30			
										~+-				++~	~~~	1 4 4
	cct															144
GIN	Pro		GTĀ	ser	ren	Arg	ьеи 40	Ard	Cys	vai	ALG	45	GIA	rne	PIO	
		35					40					#2				
++~	agt	~~~	t = t	t = c	2+~	a crt	taa	atc	ממכ	cad	act	cca	aaa	aad	aaa	192
	Ser															222
1116	50	nsp	TAT	- X -	nec	55	115	•	9		60			,	0-1	
	50					,,,					50					
tta	gag	taa	c++	aas	tta	att	aaa	acc	aat	cct	gat	aat	gga	acσ	aca	240
9	243	-99		224	LLU							25-	55-			

Leu	Glu	Trp	Leu	Gly	Leu	Ile	Lys	Thr	Asn	Pro	Asp	Gly	Gly	Thr	Thr	
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gat	tac	gcc	gcg	tct	gtg	aaa	ggc	aga	ttt	atc	atc	tca	cga	gat	gat	288
Asp	Tyr	Ala	Ala	Ser	Val	Lys	Gly	Arg	Phe	Ile	Ile	Ser	Arg	Asp	Asp	
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tca	aag	aac	tca	ctg	ttc	ctt	caa	atg	aac	agc	ctg	aaa	acc	gag	gac	336
Ser	Lys	Asn	Ser	Leu	Phe	Leu	G1n	Met	Asn	Ser	Leu	Lys	Thr	Glu	Asp	
			100					105					110			
acg	gcc	gtg	tat	tac	tgc	acc	aca	gaa	gtg	ttg	gtg	gtg	tct	gct	att	384
Thr	Ala	Val	Tyr	Tyr	Cys	Thr	Thr	Glu	Val	Leu	Val	Val	Ser	Ala	Ile	
		115					120					125				
caa	ctc	att	gga	tgt	ctg	ggg	ccc	ggg	gag	ttg	tgg	tca	ccc	gtc	tct	432
Gln	Leu	Ile	Gly	Суз	Leu	Gly	Pro	Gly	Glu	Leu	Trp	Ser	Pro	Val	Ser	
	130					135					140					
ttc	cgc	ttc	a													442
Phe	Arg	Phe														
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Va1	Leu	Ser	Gln	Val	Gln	Leu	Glu	Glu	Ser	Gly	Pro	Gly	Leu	Val	Lys	
			20					25					30			
ccc	tcg	gag	acc	ctg	tcc	ctc	acc	tgc	gct	gtg	tct	ggt	ggc	ctc	att	144
Pro	Ser	Glu	Thr	Leu	Ser	Leu	Thr	Cys	Ala	Val	Ser	Gly	Gly	Leu	Ile	
		35					40					45				
act	gga	aac	tac	tgg	aac	tgg	ctc	cgg	cag	tca	gaa	ggg	aag	gga	ctg	192
Thr	Gly	Asn	Tyr	Trp	Asn	Trp	Leu	Arg	Gln	Ser	Glu	Gly	Lys	Gly	Leu	
	50					55					60					
gag	tgg	att	ggc	cat	att	ggt	ggt	agt	agt	ggg	aac	acc	ggc	tac	aac	240
Glu	Trp	Ile	Gly	His	Ile	Gly	Gly	Ser	Ser	Gly	Asn	Thr	Gly	Tyr	Asn	
65					70					75					80	
tcc	gct	ttc	gag	agt	cgc	gtc	acc	ttg	tca	aga	gac	acg	gcc	aag	aat	288
Ser	Ala	Phe	Glu	Ser	Arg	Val	Thr	Leu	Ser	Arg	Asp	Thr	Ala	Lys	Asn	
				85					90					95		
cgg	tt¢	tcc	ctg	aaa	ctg	acc	tct	gtg	acc	gcc	gca	gat	tcg	gcc	gtc	336
Arg	Phe	Ser	Leu	Lys	Leu	Thr	Ser	Val	Thr	Ala	Ala	Asp	Ser	Ala	Val	
			100					105					110			
tat	tac	tgt	gcg	aga	tcg	ggt	ttt	acc	ggc	acc	gac	ttc	ttt	tac	tat	384
Tyr	Tyr	Cys	Ala	Arg	Ser	Gly	Phe	Thr	Gly	Thr	Asp	Phe	Phe	Tyr	Tyr	
		115					120					125				
tgg	ggc	ccg	ggg	aag	tct	tgg	tc									407
Trp	Gly	Pro	Gly	Lys	Ser	Trp										
	130					135										

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1		5				10					15		

gt	c	ctg	tcc	cag	gtt	caa	cta	cag	gag	tcg	ggc	cca	gga	ctg	atg	aag	96
Vč	<b>1</b>	Leu	Ser	Gln	Val	Gln	Leu	Gln	G1u	Ser	Gly	Pro	Gly	Leu	Met	Lys	
				20					25					30			

c	ct	tcg	gag	acc	ctg	tcc	ctc	acc	tgc	gct	gtc	tct	ggt	ggc	tcc	atc	144	
F	ro	Ser	Glu	Thr	Leu	Ser	Leu	Thr	Cys	Ala	Va1	Ser	Gly	Gly	Ser	Ile		
			35					40					45					

agc	ggt	ggt	ttt	ggc	tgg	ggc	tgg	atc	cgt	cag	tcc	ccg	ggg	aag	aaa	192
Ser	Gly	Gly	Phe	Gly	Trp	Gly	Trp	Ile	Arg	Gln	Ser	Pro	Gly	Lys	Gly	
	50					55					60					

ctg	gaa	tgg	att	gga	agt	ttc	tat	act	act	act	gga	aat	acc	ttc	tcc	240
Leu	Glu	Trp	Ile	Gly	Ser	Phe	Tyr	Thr	Thr	Thr	${\tt Gly}$	Asn	Thr	Phe	Ser	
65					70					75					80	

aac	ccc	tcc	ctc	aag	agt	cga	gtc	acc	att	tca	gcg	gac	acg	tcc	aag	288
Asn	Pro	Ser	Leu	Lys	Ser	Arg	Val	Thr	Ile	Ser	Ala	Asp	Thr	Ser	Lys	
				85					90					95		

aac	cag	ttc	tcc	ctg	aga	ctg	acc	tct	gtg	acc	gcc	gcg	gac	acg	gcc	336
Asn	Gln	Phe	Ser	Leu	Arg	Leu	Thr	Ser	Val	Thr	Ala	Ala	Asp	Thr	Ala	
			100					105					110			

gtt	tat	tac	tgt	gcg	aga	gat	ctc	tat	agc	agc	ggc	tat	aaa	ttt	tac	384
Val	Tyr	Tyr	Cys	Ala	Arg	Asp	Leu	Tyr	Ser	Ser	Gly	Tyr	Lys	Phe	Tyr	

PCT/US99/09131 WO 99/55369

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Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly 15 1

10

Ser Leu Arg Leu Ala Cys Val Gly Ser Gly Phe Ala Phe Arg Asn Thr

Arg Met His Trp Ile Arg Gln Thr Pro Gly Lys Arg Leu Glu Trp Val

45 40 35

Ala Asp Ile Lys Phe Asp Gly Ser Asp Phe Tyr Tyr Val Asp Ser Val

50 55

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Ser Leu Tyr 75 70

Leu Glu Met Asn Ser Leu Arg Pro Asp Asp Thr Ala Val Tyr Phe Cys

90 95 85

Val Arg

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Glu Val His Leu Val Gln Ser Gly Ala Gln Val Lys Arg Pro Gly Glu

1 5 10 15

Ser Leu Arg Ile Ser Cys Lys Thr Ser Gly Tyr Thr Phe Thr Asp Ser

20 25 30

Trp Ile Ser Trp Val Arg Gln Met Pro Gly Lys Gly Leu Glu Trp Met

35 40 45

Gly Asn Ile Tyr Pro Gly Asp Ser Asp Ser Arg Tyr Asn Pro Ser Phe

50 55 60

Gln Gly Arg Val Thr Ile Ser Val Asp Lys Ser Ile Ser Thr Thr Tyr

65 70 75 80

Leu Gln Trp Ser Ser Leu Lys Ala Ser Asp Thr Ala Thr Tyr Tyr Cys

85 90 95

Ala Lys

<210> 48

<211> 98

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Ala Arg

<210> 49

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<223> CDRII

<400> 49

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1 5 10 15

Thr Leu Ser Leu Thr Cys Asn Val Ser Gly Asp Ser Pro Thr Lys Ser
20 25 30

Thr Trp Asn Trp Val Arg Gln Ser Pro Gly Lys Pro Leu Glu Trp Ile

35 40 45

Gly His Val Gly Ser Gly Gly Gly Pro Val Tyr Asn Val Phe Leu 55 Thr Gly Arg Val Ser Met Ser Leu Asp Ala Ser Lys Leu Leu Ser 75 65 70 80 Leu Ala Leu Ala Ser Val Thr Ala Ala Asp Ser Ala Val Tyr Tyr Cys 85 90 95 Val Arg <210> 50 <211> 100 <212> PRT <213> Macaca cynomolgus <220> <221> DOMAIN <222> (31)...(35) <223> CDRI <221> DOMAIN <222> (50)...(68) <223> CDRII <400> 50 Asp Lys Gln Leu Val Gln Ser Gly Gly Gly Leu Val Gln Pro Gly Gly Ser Leu Arg Leu Ala Cys Val Ala Ser Gly Phe Pro Phe Ser Asp Tyr 25 30 20 Tyr Met Ser Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Leu

65 70 75 80

Leu Phe Leu Gln Met Asn Ser Leu Lys Thr Glu Asp Thr Ala Val Tyr

Tyr Cys Thr Thr

85

90

100

<210> 51

<211> 98

<212> PRT

<213> Macaca cynomolgus

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Gln Val Gln Leu Glu Glu Ser Gly Pro Gly Leu Val Lys Pro Ser Glu

1 5 10 15

Thr Leu Ser Leu Thr Cys Ala Val Ser Gly Gly Leu Ile Thr Gly Asn

20 25 30

Tyr Trp Asn Trp Leu Arg Gln Ser Glu Gly Lys Gly Leu Glu Trp Ile

35 40 45

Gly His Ile Gly Gly Ser Ser Gly Asn Thr Gly Tyr Asn Ser Ala Phe

50 55 60

Glu Ser Arg Val Thr Leu Ser Arg Asp Thr Ala Lys Asn Arg Phe Ser
65 70 75 80

Leu Lys Leu Thr Ser Val Thr Ala Ala Asp Ser Ala Val Tyr Tyr Cys

85 90 95

Ala Arg

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<400> 53

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			20					25					30			
											act					144
Leu	Ser	Thr	Ser	Val	Gly	Asp		Val	Thr	Ile	Thr		Arg	Ala	Ser	
		35					40					45				
																400
											cag					192
Gln		Ile	Asp	Thr	Glu		Ala	Trp	Tyr	GIn	Gln	гЛS	Pro	GTĀ	гÃг	
	50					55					60					
								~~~	***	200	++~	cac	200	aaa	ata	240
											ttg					240
	Pro	Thr	Leu	ьeu	70	ser	Asp	Ата	ser	75	Leu	GIII	F 4.4.T	GIY	80	
65					70					, ,					00	
tca	tet	caa	ttc	age	aac	agt.	gga	tet	gga	aca	gat	ttc	act	ctc	acc	288
											Asp					
	501			85	· · · ·		~ <i></i>		90					95		
atc	aac	agc	ctg	cag	cct	gaa	gat	att	gcg	act	tat	tac	tgt	caa	cag	336
											Tyr					
			100					105					110			
gat	aat	agt	ttt	cca	ctc	act	ttc	ggc	gga	ggg	acc	aag	gtg	gag	atc	384
Asp	Asn	Ser	Phe	Pro	Leu	Thr	Phe	Gly	Gly	Gly	Thr	Lys	Val	Glu	Ile	
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aaa	cga															390
Lys	Arg															
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<212> DNA

<213> Macaca cynomolgus

<220>

<221> CDS

<222> (1)...(384)

<400> 54

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gtg	atg	acc	cag	tct	cca	gac	tcc	ctg	gct	gtg	tct	ctg	gga	gag	agg	96
Val	Met	Thr	Gln	Ser	Pro	Asp	Ser	Leu	Ala	Val	Ser	Leu	Gly	Glu	Arg	
			20					25					30			

gtc	acc	atc	aat	tgt	aag	tcc	agc	cag	agt	ctt	tta	tac	agc	tcc	aac	144
Val	Thr	Ile	Asn	Суз	Lys	Ser	Ser	${\tt Gln}$	Ser	Leu	Leu	Tyr	Ser	Ser	Asn	
		35					40					45				

aat	aag	aac	tac	tta	gcc	tgg	tac	cag	caa	aaa	cca	gga	cag	gct	cct	192
Asn	Lys	Asn	Tyr	Leu	Ala	Trp	Tyr	Gln	Gln	Lys	Pro	gly	Gln	Ala	Pro	
	50					55					60					

caa	cta	ctc	att	tac	tgg	gca	tct	acc	cgg	gaa	tcc	ggg	gtc	cct	aat	2	40
Gln	Leu	Leu	Ile	Tyr	Trp	Ala	Ser	Thr	Arg	Glu	Ser	Gly	Val	Pro	Asn		
65					70					75					80		

cga	ttt	agt	ggc	agc	ggc	tct	ggg	aca	gat	ttc	act	ctc	acc	atc	agt	288
Arg	Phe	Ser	Gly	Ser	Gly	Ser	Gly	Thr	Asp	Phe	Thr	Leu	Thr	Ile	Ser	
				85					90					95		

ggc	ctg	cag	gct	gaa	gat	gtg	gca	gtg	tat	tac	tgt	caa	cag	tat	tat	336
Gly	Leu	Gln	Ala	Glu	Asp	Val	Ala	Val	Tyr	Tyr	Cys	Gln	Gln	Tyr	Tyr	
			100					105					110			

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<210> 55

<211> 399

<212> DNA

<213> Macaca cynomolgus

<220>

<221> CDS

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<400> 55

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Met Arg Leu Pro Ala Gln Leu Leu Gly Leu Leu Leu Cys Val Pro

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gga tcc agt ggg gat gtt gtg atg act cag tct cca ctc tcc ctg ccc 96

Gly Ser Ser Gly Asp Val Val Met Thr Gln Ser Pro Leu Ser Leu Pro

20 25 30

gtc atc cct gga cag cca gcc tcc atc tcc tgc agg tct agt caa agc 144

Val Ile Pro Gly Gln Pro Ala Ser Ile Ser Cys Arg Ser Ser Gln Ser

35 40 45

ctt gta cat agt gac ggg aaa acc tac ttg aat tgg tta caa cag aag 192
Leu Val His Ser Asp Gly Lys Thr Tyr Leu Asn Trp Leu Gln Gln Lys
50 55 60

cca ggc caa cct cca aga ctc ctg att tat cag gtt tct aac cgg cac

Pro Gly Gln Pro Pro Arg Leu Leu Ile Tyr Gln Val Ser Asn Arg His

65 70 75 80

tct ggg gtc cca gac aga ttc agc ggc agt ggg gca ggg aca gac ttc 288
Ser Gly Val Pro Asp Arg Phe Ser Gly Ser Gly Ala Gly Thr Asp Phe
85 90 95

aca	ctg	aaa	atc	agc	aga	gtg	gag	act	gag	gat	gtt	ggg	gtt	tat	tcc	336
Phr	Leu	Lys	Ile	Ser	Arg	Val	Glu	Thr	Glu	Asp	Val	Gly	Val	$\mathtt{Tyr}$	Ser	
			100					105					110			
tgc	gtg	caa	ggt	aca	cac	tgg	ccg	tgg	acg	ttc	ggc	çaa	ggg	acc	aag	384
Cys	Val	Gln	Gly	Thr	His	Trp	Pro	Trp	Thr	Phe	Gly	${\tt Gln}$	Gly	Thr	Lys	
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gtg	gac	atc	aaa	cga												399
Val	Asp	Ile	Lys	Arg												
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Met	Arg	Val	Pro	Ala	Gln	Leu	Leu	Gly	Leu	Leu	Leu	Leu	Trp	Leu	Pro	
1				5					10					15		
ggt	gcc	ata	tgt	gac	att	cag	atg	tcc	cag	tct	cca	tcc	tcc	ctg	tct	96
Gly	Ala	Ile	Cys	Asp	Ile	Gln	Met	Ser	Gln	Ser	Pro	Ser	Ser	Leu	Ser	
			20					25					30			
gct	tct	gtg	gga	gac	aga	gtc	acc	atc	acc	tgc	cgg	gca	agt	cag	ggc	144
Ala	Ser	Val	Gly	Asp	Arg	Val	Thr	Ile	Thr	Cys	Arg	Ala	Ser	Gln	Gly	
		35					40					45				

57

Ile Thr Asn Tyr Leu Asn Trp Tyr Gln Gln Lys Pro Gly Lys Ala Pro 50 55 60 240 aac ctc ctg atc tat tat gca act cgt ttg gcg agc ggg gtc cca tca Asn Leu Leu Ile Tyr Tyr Ala Thr Arg Leu Ala Ser Gly Val Pro Ser 65 70 75 80 agg ttc agc ggc agt gga tct ggg tcg gag tac agt ctc gcc atc agc 288 Arg Phe Ser Gly Ser Gly Ser Glu Tyr Ser Leu Ala Ile Ser 90 95 85 age etg cag eet gaa gat ttt gea ace tat tte tgt caa cag ggt tat 336 Ser Leu Gln Pro Glu Asp Phe Ala Thr Tyr Phe Cys Gln Gln Gly Tyr 100 105 110 384 agg gcc ccc tac act ttt ggc cag ggg acc aca gtg gag atc aaa cga Arg Ala Pro Tyr Thr Phe Gly Gln Gly Thr Thr Val Glu Ile Lys Arg 115 120 125 <210> 57 <211> 390 <212> DNA <213> Macaca cynomolgus <220> <221> CDS <222> (1)...(390) <400> 57 48 atg gac atg agg gtc ccc gct cag ctc ctg ggg ctc ctg ctg ctc tgg Met Asp Met Arg Val Pro Ala Gln Leu Leu Gly Leu Leu Leu Trp 1 10 15 ctc cta ggt gcc aga tgt gac atc cag atg acc cag tct cct tct tcc 96 Leu Leu Gly Ala Arg Cys Asp Ile Gln Met Thr Gln Ser Pro Ser Ser 30 20 25

ttg	tct	gca	tct	gta	gga	gac	aga	gtc	acc	atc	act	tgc	caa	gcc	agt	1	44
Leu	Ser	Ala	Ser	Val	Gly	Asp	Arg	Va1	Thr	Ile	Thr	Cys	Gln	Ala	Ser		
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cag	ggt	att	agc	aac	tgg	tta	gcc	tgg	tat	cag	cag	aaa	ccg	aaa	aaa	1	.92
Gln	G1y	Ile	Ser	Asn	Trp	Leu	Ala	Trp	Tyr	Gln	Gln	Lys	Pro	Gly	Lys		
	50					55					60						
						tat										2	40
Ala	Pro	Lys	Leu	Leu	Ile	Tyr	Ala	Ala	Ser	Thr	Phe	Gln	Ser	Gly	Val		
65					70					75					80		
						agt										2	88
Pro	Ser	Arg	Phe	Ser	Gly	Ser	Gly	Ser		Thr	Glu	Phe	Thr		Thr		
				85					90					95			
																_	
						gaa										3	336
Ile	Ser	Ser		Gln	Pro	Glu	Asp		Ala	Thr	Tyr	Tyr		GIn	GIn		
			100					105					110				
							L +			~~~	200	~	~+ <i>~</i>	a a a	atc	2	384
						act										-	,04
туr	ASI		TYT	Pro	Leu	Thr	120	GTĀ	Слу	Gry	1111	125	Val	Ora	110		
		115					120										
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	Arg																
_,_	130																
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1				5					10					15		
ctc	cca	ggt	gcc	aga	ggt	gac	atc	cag	atg	acc	cag	tct	cca	ccc	tcc	96
Leu	Pro	Gly	Ala	Arg	Gly	Asp	Ile	Gln	Met	Thr	Gln	Ser	Pro	Pro	Ser	
			20					25					30			
ctg	tet	gcg	tct	gtt	ggg	gac	act	gtc	agt	ctt	act	tgt	cgg	gca	agt	144
Leu	Ser	Ala	Ser	Val	Gly	Asp	Thr	Val	Ser	Leu	Thr	Cys	Arg	Ala	Ser	
		35					40					45				
cag	cct	att	ggc	agt	aat	tta	aat	tgg	ttc	cag	caa	aaa	cct	ggg	agc	192
	Pro															
	50		-			55					60					
ccc	ccc	aga	ctc	cta	atc	tac	ctt	gcg	acc	gcc	ttg	caa	cgt	ggg	atc	240
	Pro	-														
65		5			70					75				_	80	
cca	tca	agg	ttt	agc	acc	act	aaa	tct	caa	acc	aat	ttc	act	ctc	acg	288
	Ser															
	501	111 9	1110	85			1		90					95		
									-							
atc	acc	aac	cta	car	cct	nan	cat	ttc	gca	act	tac	ctc	tat	cta	caa	336
	Thr															
116	1111	GTĀ	100	GIII	FIO	GIU	nup	105	1114		*1*	2504	110	204	<b>411</b>	
			100					103					110			
~~+	act		+		++~	-a-		aac	ccc	aaa	202	ne e	ata	a=+	atc	384
	Thr															204
nis	THE		TYT	PTO	rne	THE		GTĀ	F10	GTĀ	TIIT		val	vsb	116	
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aag	cga															390

Lys Arg

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<210> 59

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<212> PRT

<213> Macaca cynomolgus

<220>

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<222> (50)...(56)

<223> CDRII

<400> 59

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Asp Thr Val Thr Ile Thr Cys Arg Ala Ser Gln Gly Ile Asp Thr Glu 20 25 30

Leu Ala Trp Tyr Gln Gln Lys Pro Gly Lys Ala Pro Thr Leu Leu Ile 35 40 45

Ser Asp Ala Ser Arg Leu Gln Thr Gly Val Ser Ser Arg Phe Ser Gly
50 55 60

Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile Asn Ser Leu Gln Pro 65 70 75 80

Glu Asp Ile Ala Thr Tyr Tyr Cys

85

<210> 60

<211> 94

<212> PRT

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10

Glu Arg Val Thr Ile Asn Cys Lys Ser Ser Gln Ser Leu Leu Tyr Ser 25

Ser Asn Asn Lys Asn Tyr Leu Ala Trp Tyr Gln Gln Lys Pro Gly Gln

45 40 35

Ala Pro Gln Leu Leu Ile Tyr Trp Ala Ser Thr Arg Glu Ser Gly Val

55 60

Pro Asn Arg Phe Ser Gly Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr 75 65 70 80

Ile Ser Gly Leu Gln Ala Glu Asp Val Ala Val Tyr Tyr Cys 90

<210> 61

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<213> Macaca cynomolgus

85

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<221> DOMAIN

<222> (24)...(39)

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<221> DOMAIN

<222> (54)...(61)

<223> CDRII

<400> 61

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<210> 62

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<220>

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<223> CDRII

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Asp Arg Val Thr Ile Thr Cys Arg Ala Ser Gln Gly Ile Thr Asn Tyr
20 25 30

Leu Asn Trp Tyr Gln Gln Lys Pro Gly Lys Ala Pro Asn Leu Leu Ile
35 40 45

Tyr Tyr Ala Thr Arg Leu Ala Ser Gly Val Pro Ser Arg Phe Ser Gly 50 55 60

Ser Gly Ser Gly Ser Glu Tyr Ser Leu Ala Ile Ser Ser Leu Gln Pro

65 70 75 80

Glu Asp Phe Ala Thr Tyr Phe Cys

85

<210> 63

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1 5 10 15

Asp Arg Val Thr Ile Thr Cys Gln Ala Ser Gln Gly Ile Ser Asn Trp
20 25 30

Leu Ala Trp Tyr Gln Gln Lys Pro Gly Lys Ala Pro Lys Leu Leu Ile 35 40 45

Tyr Ala Ala Ser Thr Phe Gln Ser Gly Val Pro Ser Arg Phe Ser Gly
50 55 60

Ser Gly Ser Gly Thr Glu Phe Thr Leu Thr Ile Ser Ser Leu Gln Pro 65 70 75 80

Glu Asp Phe Ala Thr Tyr Tyr Cys

85

<210> 64

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10

15

Asp Thr Val Leu Thr Gln Ser Pro Ala Leu Ala Val Pro Pro Gly Glu

5

1

agg	gtt	acc	gtc	tcc	tgt	agg	gcc	agt	gaa	agt	gtc	agt	aca	ttt	ttg	96
Arg	Val	Thr	Val	Ser	Cys	Arg	Ala	Ser	Glu	Ser	Val	Ser	Thr	Phe	Leu	
			20					25					30			
cac	tgg	tat	caa	cag	aaa	cca	gga	cat	caa	CCC	aaa	ctc	ctc	atc	tat	144
His	Trp	Tyr	Gln	Gln	Lys	Pro	Gly	His	Gln	Pro	Lys	Leu	Leu	Ile	Tyr	
		35					40					45				
cta	gcc	tca	aaa	cta	gaa	tct	aaa	gtc	cct	gcc	agg	ttc	agt	ggc	ggt	192
Leu	Ala	Ser	Lys	Leu	Glu	Ser	Gly	Val	Pro	Ala	Arg	Phe	Ser	Gly	Gly	
	50					55					60					
ggg	tct	ggg	aca	gac	ttc	acc	ctc	acc	att	gat	cct	gtg	gag	gct	gat	240
Gly	Ser	Gly	Thr	Asp	Phe	Thr	Leu	Thr	Ile	Asp	Pro	Val	Glu	Ala	Asp	
65					70					75					80	
gac	act	gct	acc	tat	tac	tgt	cag	cag	acc	tgg	aat	gat	cct	cgg	acg	288
Asp	Thr	Ala	Thr	Tyr	Tyr	Cys	Gln	Gln	Thr	Trp	Asn	Asp	Pro	Arg	Thr	
				85					90					95		
				0,5												
		gga		acc					aaa					gca		336
		gga Gly		acc					aaa				Ala	gca		336
				acc					aaa					gca		336
			Gly	acc				Leu	aaa				Ala	gca		
Phe	Gly		Gly 100	acc Thr	Lys	Leu	Glu	Leu	aaa				Ala	gca		336 360
Phe	Gly gta	Gly	Gly 100 atc	acc Thr	Lys cca	Leu cca	<b>G</b> lu	Leu	aaa				Ala	gca		
Phe	Gly gta	Gly	Gly 100 atc	acc Thr	Lys cca	Leu cca	<b>G</b> lu	Leu	aaa				Ala	gca		
Phe	Gly gta	Gly tct Ser	Gly 100 atc	acc Thr	Lys cca	Leu cca	Glu tcc Ser	Leu	aaa				Ala	gca		
Phe	Gly gta Val	tct Ser 115	Gly 100 atc Ile	acc Thr	Lys cca	Leu cca	Glu tcc Ser	Leu	aaa				Ala	gca		
Phe	Gly gta Val	tct Ser 115	Gly 100 atc Ile	acc Thr	Lys cca	Leu cca	Glu tcc Ser	Leu	aaa				Ala	gca		
Phe	gta Val	tct Ser 115 210>	Gly 100 atc Ile 66 360	acc Thr	Lys cca	Leu cca	Glu tcc Ser	Leu	aaa				Ala	gca		
Phe	gta Val	Gly  tct Ser 115 210> 211> 212>	Gly 100 atc Ile 66 360 DNA	acc Thr	Lys cca	Leu cca	Glu tcc Ser	Leu	aaa				Ala	gca		
Phe	gta Val	tct Ser 115 210>	Gly 100 atc Ile 66 360 DNA	acc Thr	Lys cca	Leu cca	Glu tcc Ser	Leu	aaa				Ala	gca		
Phe	gta Val	cly tct ser 115 210> 211> 212> 213>	Gly 100 atc Ile 66 360 DNA	acc Thr	Lys cca	Leu cca	Glu tcc Ser	Leu	aaa				Ala	gca		
Phe	gta Val	Gly  tct Ser 115 210> 211> 212>	Gly 100 atc Ile 66 360 DNA Rat	acc Thr	Lys cca	Leu cca	Glu tcc Ser	Leu	aaa				Ala	gca		

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Glu	Val	Gln	Leu	Gln	Gln	Ser	Gly	Pro	Glu	Val	Gly	Arg	Pro	Gly	Ser		
1				5					10					15			
tca	gte	aag	att	tat	tgc	aag	gct	tct	ggc	tac	acc	ttt	aca	gat	tac	<u>.</u>	96
Ser	Val	Lys	Ile	Ser	Cys	Lys	Ala	Ser	Gly	Tyr	Thr	Phe	Thr	Asp	Tyr		
			20					25					30				
gtt	ttg	aat	tgg	gtg	aag	cag	agt	cct	gga	cag	gga	ctg	gaa	tgg	ata	14	44
Val	Leu	Asn	Trp	Va1	Lys	Gln	Ser	Pro	Gly	Gln	Gly	Leu	Glu	Trp	Ile		
		35					40					45					
gga	tgg	att	gat	cct	gac	tat	ggt	act	act	gat	tat	gct	gag	aag	ttc	19	92
Gly	Trp	Ile	Asp	Pro	Asp	Tyr	Gly	Thr	Thr	Asp	Tyr	Ala	Glu	Lys	Phe		
	50					55					60						
aaa	aag	aag	gcc	aca	ctg	act	gca	gat	aca	tcc	tcc	agc	aca	gcc	tac	24	40
Lys	Lys	Lys	Ala	Thr	Leu	Thr	Ala	Asp	Thr	Ser	Ser	Ser	Thr	Ala	Tyr		
65					70					75					80		
atc	cag	ctt	agc	agc	ctg	aca	tct	gag	gac	aca	gcc	acc	tat	ttt	tgt	28	88
Ile	Gln	Leu	Ser	Ser	Leu	Thr	Ser	Glu	Asp	Thr	Ala	Thr	Tyr	Phe	Суз		
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gct	aga	tct	agg	aat	tac	gga	gga	tat	att	aat	tac	tgg	ggc	caa	gga	3:	36
Ala	Arg	Ser	Arg	Asn	Tyr	Gly	Gly	Tyr	Ile	Asn	Tyr	Trp	Gly	Gln	Gly		
			100					105					110				
gtc	atg	gtc	aca	gtc	tcc	tca	gct									3 (	60
Val	Met	Val	Thr	Val	Ser	Ser	Ala										
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<212> PRT

<213> Pan troglodytes

<400> 67

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20 25 30

Leu Asn Trp Tyr Gln Gln Lys Pro Gly Lys Ala Pro Lys Leu Leu Ile 35 40 45

Phe Asp Ala Ser Ile Leu Gln Ser Gly Val Pro Ser Arg Phe Ser Gly

Ser Gly Ser Gly Thr Asp Phe Ser Leu Thr Ile Arg Ser Leu Gln Pro 65 70 75 80

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Tyr Asn Phe Gly Gln Gly Thr Lys Leu Glu Ile Lys Arg 100 105

<210> 68

<211> 108

<212> PRT

<213> Artificial Sequence

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<223> rat/chimpanzee sequence

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20 25 30

Leu His Trp Tyr Gln Gln Lys Pro Gly Lys Ala Pro Lys Leu Leu Ile 35 40 45

Tyr Leu Ala Ser Lys Leu Glu Ser Gly Val Pro Ala Arg Phe Ser Gly 50 55 60

Ser Gly Ser Gly Thr Asp Phe Ser Leu Thr Ile Arg Ser Leu Gln Pro

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Val	Leu	Asn	Trp	Va1	Lys	Gln	Ala	Pro	Gly	Lys	Gly	Leu	Glu	Trp	Ile		
		35					40					45					
Gly	Trp	Ile	Asp	Pro	Asp	Tyr	Gly	Thr	Thr	Asp	Tyr	Ala	Glu	Lys	Phe		
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Ala	Arg	Ser	Arg	Asn	Tyr	Gly	Gly	Tyr	Ile	Asn	Tyr	Trp	Gly	Gln	Gly		
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1				5					10					15			
tca	gtg	aag	ata	tcc	tgc	aag	gct	act	ggc	tac	aca	ttc	agt	agc	tac	!	96
Ser	Val	Lys	Ile	Ser	Cys	Lys	Ala	Thr	Gly	Tyr	Thr	Phe	Ser	Ser	Tyr		
			20					25					30				

144

tgg ata gag tgg gta aag cag agg cct gga cat ggc ctt gag tgg att

Trp Ile Glu Trp Val Lys Gln Arg Pro Gly His Gly Leu Glu Trp Ile 45 35 40 192 gga gag att tta cct aga agt ggt aat act aac tac aat gag aag ttc Gly Glu Ile Leu Pro Arg Ser Gly Asn Thr Asn Tyr Asn Glu Lys Phe 50 55 60 aag ggc aag gcc aca ttc act gca gaa aca tcc tcc aac aca gcc tac 240 Lys Gly Lys Ala Thr Phe Thr Ala Glu Thr Ser Ser Asn Thr Ala Tyr 65 70 75 80 atg caa ctc agc agc ctg aca cct gag gac tct gcc gtc tat tac tgt 288 Met Gln Leu Ser Ser Leu Thr Pro Glu Asp Ser Ala Val Tyr Tyr Cys 90 95 85 tca agt cgc ggc gtc agg ggc tct atg gac tac tgg ggt caa gga acc 336 Ser Ser Arg Gly Val Arg Gly Ser Met Asp Tyr Trp Gly Gln Gly Thr 105 110 100 354 tca gtc acc gtc tcc tca Ser Val Thr Val Ser Ser 115 <210> 72 <211> 324 <212> DNA <213> Murine <220> <221> CDS <222> (1)...(324) <400> 72 48 gat att cag atg acc cag act aca tcc tcc ctg tct gcc tct ctg gga Asp Ile Gln Met Thr Gln Thr Thr Ser Ser Leu Ser Ala Ser Leu Gly 1 10 15

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														aat			96
Asp	Arg	Val		TTE	Thr	Cys	Arg		ser	GIII	Asp	TIG		Asn	File		
			20					25					30				
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														ctg			144
ьец	ASI		туг	GIN	GIII	гуѕ		ASD	GTÅ	1111	vaı	цу.s 45	ьеи	Leu	TIE		
		35					40					4.0					
<b>+</b>							t a a	~~~	ato	cca	toa	200	++~	a ort	aaa		192
														agt Ser			192
TAL		Thr	ser	Thr	ьeu		ser	GIY	val	PIG	60	ALG	FILE	Set	GIA		
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a.c.t	aaa	+ a +	~~~	202	œ= t	+ a +	tet	ctc	200	att	acc	aac	cta	gag	Caa		240
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65	GIY	per	GTĀ	1111	70	* Y L	DCI	ДСС		75	001	*1011	u	014	80		
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caa	gat	a++	acc	act	tac	+++	tac	caa	caq	aat	aat	acq	ctt	cct	taa		288
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GIU	изр	116	ALG	85	IYI	1110	Cys	O.1.1	90	013	11011			95			
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acq	ttc	aat	gga	aac	acc	aac	cta	gaa	atc	aaa	caa						324
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73

105

Met Glu Leu Thr Ser Leu Arg Ser Glu Asp Thr Ala Val Tyr Tyr Cys

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Ser Val Lys Leu Ser Cys Lys Ala Ser Gly Ser Thr Phe Thr Ser Tyr
20 25 30

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Trp Met His Trp Val Lys Gln Arg Pro Gly Arg Gly Leu Glu Trp Ile

35
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45

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Gly Arg Ile Asp Pro Asn Ser Gly Gly Thr Lys Asp Asn Glu Lys Phe
50 55 60

aag agc aag gcc aca ctg act gta gac aaa ccc tcc agc aca gcc tac 240

Lys Ser Lys Ala Thr Leu Thr Val Asp Lys Pro Ser Ser Thr Ala Tyr

65 70 75 80

atg cag ctc agc agc ctg aca tct gag gac tct gcg gtc tat tat tgt 288

Met Gln Leu Ser Ser Leu Thr Ser Glu Asp Ser Ala Val Tyr Tyr Cys

85 90 95

gca aga gag acc tac tat gat tcc tcg ttt gct tac tgg ggc caa ggg 336

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PCT/US99/09131 WO 99/55369

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Trp Met His Trp Val Lys Gln Ala Pro Gly Gln Gly Leu Glu Trp Ile

Gly Arg Ile Asp Pro Asn Ser Gly Gly Thr Lys Asp Asn Glu Lys Phe
50 55 60

Lys Ser Lys Ala Thr Leu Asn Val Asp Lys Ser Thr Asn Ile Ala Tyr 65 70 75 80

Met Glu Leu Thr Ser Leu Arg Ser Glu Asp Thr Ala Val Tyr Tyr Cys
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Thr Met Val Thr Val Ser

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Trp Met His Trp Val Lys Gln Ala Pro Gly Gln Gly Leu Glu Trp Ile

35 40 45 Gly Arg Ile Asp Pro Asn Ser Gly Gly Thr Lys Asp Asn Glu Lys Phe 55 Lys Ser Lys Ala Thr Leu Thr Val Asp Lys Ser Thr Ser Thr Ala Tyr 70 75 80 Met Glu Leu Ser Ser Leu Arg Ser Glu Asp Thr Ala Val Tyr Tyr Cys 90 85 Ala Arg Glu Thr Tyr Tyr Asp Ser Ser Phe Ala Tyr Trp Gly Gln Gly 100 105 110 Thr Met Val Thr Val Ser Ala 115 <210> 80 <211> 102 <212> PRT <213> Artificial Sequence <220> <223> murine/human sequence <400> 80 Asp Ile Val Met Thr Gln Ser Pro Ser Ser Leu Ser Ala Ser Val Gly 15 10 Asp Arg Val Thr Ile Thr Cys Lys Ala Ser Gln Asn Val Gly Thr Asn 25 20 Val Ala Trp Tyr Gln Gln Lys Pro Gly Lys Ala Pro Lys Ala Leu Ile 40 Tyr Ser Ala Ser Tyr Arg Tyr Ser Gly Val Pro Asp Arg Phe Ser Gly 55 60 50 Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile Ser Ser Leu Gln Pro 70 75 Glu Asp Phe Ala Thr Tyr Tyr Cys Gln Gln Tyr Asn Ser Tyr Pro Leu 95 90 Thr Phe Gly Gly Gly Thr 100 <210> 81

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<400> 97

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## INTERNATIONAL SEARCH REPORT

International application No. PCT/US99/09131

A. CLASSIFICATION OF SUBJECT MATTER  IPC(6): A61K 39/395  US CL: 530/387.3; 424/133.1  According to International Patent Classification (IPC) or to both national classification and IPC									
B. FIELDS SEARCHED									
Minimum documentation searched (classification system	followed by classification symbols)								
U.S. : 530/387.3; 424/133.1									
Documentation searched other than minimum documentati	on to the extent that such documents are included in the fields searched								
none									
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)									
APS, Medline, Biosis search terms: immunoglobulin, antibody, framework regions, CDR grafted, humanized, primatized									
C. DOCUMENTS CONSIDERED TO BE RELEVANT									
Category* Citation of document, with indication, v	where appropriate, of the relevant passages Relevant to claim No.								
receptor modulation without ma Chimpanzees: In vitro and in viv CE9.1) to human CD4.	ized MAb to Human CD4 causes rked reduction in CD4+ T cells in o characterization of a MAb (IDEC-Clinical Immunology and Vol. 84, No. 1, pages 73-84, see								
Further documents are listed in the continuation of	of Box C. See patent family annex.								
Special categories of cited documents:	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand								
*A* document defining the general state of the art which is not considered the principle or theory underlying the invention to be of particular relevance									
*B* earlier document published on or after the international filis									
*L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other sys									
special reason (as specified)  "O"  document referring to an oral disclosure, use, exhibition or other means  "O"  document referring to an oral disclosure, use, exhibition or other means  "O"  special reason (as specified)  considered to involve an inventive step when the document combined with one or more other auch documents, such comb being obvious to a person skilled in the art									
*P* document published prior to the international filing date but later than *A* document member of the same patent family the priority date claimed									
Date of the actual completion of the international search  Date of mailing of the international search report									
26 JULY 1999 18 AUG 1999									
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231	JULIE BURKÉ Naurence For								
Ferrimile No. (703) 305-3230	Telephone No. (703) 308-0196								

## INTERNATIONAL SEARCH REPORT

International application No. PCT/US99/09131

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)						
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:						
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:						
2. X Claims Nos.: 20-31 because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:  the claim contain specific sequence identification numbers however the application has not complied with the sequence requirements.						
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).						
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)						
This International Searching Authority found multiple inventions in this international application, as follows:						
1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.						
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.						
3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:						
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:						
Remark on Protest  The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.						